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64 CD 25 binding molecules.

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#### Description

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This Invention relates to immunosuppression and more particularly provides monoclonal antibodies and other binding molecules against the CD25 antigen.

In organ transplant surgery, particularly kidney, liver, heart, lung and bone marrow transplant surgery, it is necessary to suppress the immune system of the graft recipient to minimise the likelihood of graft rejection after surgery. Various immunosuppressive drugs have been proposed for this purpose but their use has to be carefully controlled since, in addition to undesirable side-effects arising from the use of certain immunosuppressive agents, there is also the difficulty that the immunosuppressive action makes the graft recipient particularly susceptible to infection by bacteria and viruses that would be controlled by a normal immune system. Immunosuppressive agents that have been used successfully in clinical practice include steroids, azathioprine and cyclosporin A. It is necessary in clinical practice to attempt to balance the degree of immunosuppression necessary to prevent or treat graft rejection episodes with the retention of a certain amount of the recipient's immune system to combat other infectious agents and, at the same time, to keep any possible undesirable side-effects under control.

In addition to the use of immunosuppressive drugs, attention has also focused upon the use of certain monoclonal antibodies (MAbs) to suppress immune reactions, in particular, attention has been paid to monoclonal antibodies that recognise various surface antigens of T-cells. Here too, problems have been encountered in clinical practice, namely that prior art antibodies were either too powerful or not sufficiently effective, and sometimes caused severe side effects such as high fever.

These MAb's are generally designated by a CD (Cluster Determination) number assigned by successive Leucocyte Typing Workshops. Although a term such as CD3 is now frequently applied to the cell surface antigen, and a MAb to this antigen is often described as "anti-CD3", in the following description terms such as CD3, CD25 etc. will be applied to MAb's and the corresponding cell surface antigens will be described as "CD3 antigen" etc.

In particular, monoclonal antibodies to membrane antigens present on all T-cells (also called pan T-cell antigens) such as the CD3 antigen are very potent antibodies in that they have an overall suppressive activity on the immune system. Therefore, the human body may be deprived of the immediate immune response usually mediated by the memory T-cells once an infection occurs. This is certainly not desirable when attempting to prevent rather than to cure graft rejection episodes. A treatment suitable for use in prophylaxis should be essentially selective, i.e. the pool of memory T-cells should be kept intact while the category of T-cells (activated T-cells) which could be directly involved in a rejection event should be inactivated.

This desirable goal may be achieved using antibodies to activated T-cells. These T-cells are characterised by the presence of the high affinity IL-2 receptor on their membrane surface. The high affinity IL-2 receptor is composed of at least two different polypeptide chains, an  $\alpha$ -chain also known as the CD25 antigen, and a  $\beta$ -chain. Resting T-cells do not express this high affinity receptor but low and intermediate affinity receptors which consist of  $\alpha$ - or  $\beta$ -chain homodimers. A CD25 antibody which interferes with the binding of IL-2 to its high affinity receptor and therefore selectively suppresses the immune response, is an antibody of choice for the prophylaxis of graft rejection episodes.

Natural immunoglobulins or antibodies comprise a generally Y-shaped multimeric molecule having an antigen-binding site at the end of each upper arm. The remainder of the structure, in particular the stem of the Y mediates effector functions associated with the immunoglobulins. The general structure of an antibody of the IgG class is shown schematically in Figure 1A. Both heavy and light chains comprise a variable domain and a constant part. An antigen binding site consists of the variable domain of a heavy chain associated with the variable domain of a light chain. The variable domains of the heavy and light chains have the same general structure which is illustrated in Figure 1B.

More particularly, the antigen binding characteristics of an antibody are essentially determined by 3 specific regions in the variable domain of the heavy and light chains which are called hypervariable regions or complementarity determining regions (CDRs). As shown in Figure 1B, these 3 hypervariable regions alternate with 4 framework regions, (FRs) whose sequences are relatively conserved and which are not directly involved in binding. The CDRs form loops and are held in close proximity by the framework regions which largely adopt a  $\beta$ -sheet conformation. The CDRs of a heavy chain together with the CDRs of the associated light chain essentially constitute the antigen binding site of the antibody molecule.

The determination as to what constitutes an FR or a CDR region is usually made by comparing the amino acid sequence of a number of antibodies raised in the same species. The general rules for identifying the CDR and FR regions are given in Tabl I.

Furthermore, it has been recently found that the contribution made by a light chain variable domain to the energetics of binding is small compared to that made by the associated heavy chain variable domain and that

isolated heavy chain variable domains have an antigen binding activity on their own. Such molecules are now commonly ref rred to as single domain antibodies.

Several murine CD25 MAbs already exist and include 33B3-1 (Immunotech-Merieux), BDαIL-2R (Becton-Dickinson), 2C8 (Amersham), Campath 6 (MRC, Cambridge) and ATH207(free University, Berlin). However, it has now been found that a novel mouse CD25 antibody of the IgG2a isotype, hereinafter called RFT5-IgG2a, has better properties than the CD25 antibodies of the prior art especially with regard to binding affinity, and that it is possible to construct other CD25 binding molecules having the same hypervariable regions as RFT5-IgG2a.

Accordingly, the invention provides a CD25 binding molecule which comprises at least one antigen binding site comprising at least one domain which comprises in sequence, the hypervariable regions CDR1, CDR2 and CDR3; said CDR1 having the amino acid sequence Arg-Tyr-Trp-Met-His, said CDR2 having the amino acid sequence Ala-lle-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly, and said CDR3 having the amino acid sequence Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe; and direct equivalents thereof.

In a first aspect of the invention, the CD25 binding molecule comprises a single antigen binding site comprising a single domain.

In a second aspect of the invention, the CD25 binding molecule comprises at least one antigen binding site comprising:

a) a first domain comprising in sequence the hypervariable regions CDR1, CDR2 and CDR3; said CDR1 having the amino acid sequence Arg-Tyr-Trp-Met-His, said CDR2 having the amino acid sequence Ala-Ile-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly, and said CDR3 having the amino acid sequence Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe and,

b) a second domain comprising in sequence the hypervariable regions CDR1', CDR2' and CDR3', said CDR1' having the amino acid sequence Ser-Ala-Ser-Ser-Ser-Ile-Ser-Tyr-Met-Gln, said CDR2' having the amino acid sequence Asp-Thr-Ser-Lys-Leu-Ala-Ser, and said CDR3' having the amino acid sequence His-Gln-Arg-Ser-Ser-Tyr-Thr;

and direct equivalents thereof.

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Unless otherwise indicated, any polypeptide chain is hereinafter described as having an amino acid sequence starting at the N-terminal extremity and ending at the C-terminal extremity.

When the antigen binding site comprises both the first and second domains, these may be located on the same polypeptide molecule or, preferably, each domain may be on a different chain, the first domain being part of an immunoglobulin heavy chain or fragment thereof and the second domain being part of an immunoglobulin light chain or fragment thereof.

By "CD25 binding molecule" is meant any molecule capable of binding to the CD25 antigen either alone or associated with other molecules to form high affinity IL-2 receptors. The binding reaction may be shown by standard methods (qualitative assays) including, for example, a bioassay for determining the inhibition of IL-2 binding to its receptor or any kind of binding assays, with reference to a negative control test in which an antibody of unrelated specificity, e.g. an anti-lysosyme antibody, is used. Advantageously, the binding of the molecule of the invention to the CD25 antigen may be shown in a competitive binding assay using the AHT207, BDαIL-2-R or 33B3-1 antibody as competitors. Preferably, the AHT207 or BDαIL-2-R antibody will be chosen as competitors. A particular example of a binding assay is given below.

Human peripheral blood mononuclear cells (HPBM) are grown in culture medium RPMI 1640 supplemented with 2mM L-glutamine, 100 units/ml penicillin, 100 μg/ml streptomycin, 25 mM sodium bicarbonate and 10% fetal calf serum (FCS). 1 μg/ml phytohemagglutinin (PHA) is used to stimulate HPBM. After 3 days, the blasts are resuspended at a concentration of 3.10% ml in phosphate buffered saline supplemented with 2% bovine serum albumin (BSA) and 2% azide. 50 μl samples of this suspension are incubated for 10 mn, at 20°C, under non-capping conditions, with graded concentrations of a blocking antibody (competitor) from 1 to 100 μg/ml. Then 1 μg/ml of biotinylated antibody of the invention is added to the cells and the incubation is continued for 10 min. Cells are washed and further incubated for 10 min with fluorescein-labelled streptavidin. Cells are again washed, fixed with formalin and analysed with a fluoro- cytometer which detects the binding of the biotinylated antibody. In parallel, an experiment is carried out with a biotinylated antibody of an unrelated specificity, as a negative control.

Examples of antigen binding molecules include antibodies as produced by B-cells or hybridomas and chimeric or humanized antibodies or any fragment thereof, e.g. F(ab')<sub>2</sub> and Fab fragments, as well as single chain or single domain antibodies.

A single chain antibody consists of the variable domains of an antibody heavy and light chains covalently bound by a peptide linker usually consisting of from 10 to 30 amino acids, preferably from 15 to 25 amino acids. The refore, such a structure does not include the constant part of the heavy and light chains and it is believed that the small peptide spacer should be less antigenic than a whole constant part. By "chimeric antibody" is

meant an antibody in which the constant regions of heavy or light chains or both are of human origin while the variable domains of both heavy and light chains are of non-human (e.g. murine) origin. By "humanized antibody" is meant an antibody in which the hypervariable regions (CDRs) are of non-human (e.g. murine) origin, while all or substantially all the other parts of the immunoglobuline.g. the constant regions and the highly conserved parts of the variable domains, i.e. the framework regions, are of human origin. A humanized antibody may however retain a few amino acids of the murine sequence in the parts of the framework regions adjacent to the hypervariable regions.

Hypervariable regions may be associated with any kind of framework regions, preferably of murine or human origin. Suitable framework regions are described in "Sequences of proteins of immunological interest", Kabat E.A. et al, US department of health and human services, Public health service, National Institute of Health. However, the preferred heavy chain framework is that of RFT5-IgG2a, which is shown in Seq. Id. No. 1. It consists in sequence of FR1, FR2, FR3 and FR4 regions. In a similar manner, Seq. Id. No. 2 shows the preferred RFT5-IgG2a light chain framework which consists, in sequence, of FR1', FR2', FR3' and FR4' regions.

Accordingly, the invention also provides a CD25 binding molecule which comprises at least one antigen binding site comprising either a first domain having an amino acid sequence substantially identical to that shown in Seq. Id. No. 1 starting with amino acid at position 1 and ending with amino acid at position 117 or a first domain as described above and a second domain having an amino acid sequence substantially identical to that shown in Seq. Id. No. 2, starting with amino acid at position 1 and ending with amino acid at position 104.

Monoclonal antibodies raised against a protein naturally found in all humans must necessarily be developed in a non-human system e.g. in mice. As a direct consequence of this, a xenogenic antibody as produced by a hybridoma, when administered to humans, elicits an undesirable immune response which is predominantly mediated by the constant part of the xenogenic immunoglobulin. This clearly limits the use of such antibodies as they cannot be administered over a prolonged period of time. Therefore it is particularly preferred to use single chain, single domain, chimeric or humanized antibodies which are not likely to elicit a substantial allogenic response when administered to humans.

In view of the foregoing, a more preferred CD25 binding molecule of the invention is selected from a chimeric anti-CD25 antibody which comprises at least

a) one immunoglobulin heavy chain or fragment thereof which comprises (i) a variable domain comprising in sequence the hypervariable regions CDR1, CDR2 and CDR3 and (ii) the constant part or fragment thereof of a human heavy chain; said CDR1 having the amino acid sequence Arg-Tyr-Trp-Met-His, said CDR2 having the amino acid sequence Ala-Ile-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly, and said CDR3 having the amino acid sequence Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe and

b) one immunoglobulin light chain or fragment thereof which comprises (i) a variable domain comprising in sequence the hypervariable regions CDR1', CDR2' and CDR3' and (ii) the constant part or fragment thereof of a human light chain; said CDR1' having the amino acid sequence Ser-Ala-Ser-Ser-Ser-Ile-Ser-Tyr-Met-Gln, said CDR2' having the amino acid sequence Asp-Thr-Ser-Lys-Leu-Ala-Ser, and said CDR3' having the amino acid sequence His-Gln-Arg-Ser-Ser-Tyr-Thr;

and direct equivalents thereof.

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Alternatively, a CD25 binding molecule of the invention may be selected from a single chain binding molecule which comprises an antigen binding site comprising

- a) a first domain comprising in sequence the hypervariable regions CDR1, CDR2 and CDR3, said hypervariable regions having the amino acid sequences as shown in Seq. Id. No. 1,
- b) A second domain comprising in sequence the hypervariable regions CDR1', CDR2' and CDR3', said hypervariable regions having the amino acid sequences as shown in Seq. Id. No. 2 and
- c) a peptide linker which is bound either to the N-terminal extremity of the first domain and to the C-terminal extremity of the second domain or to the C-terminal extremity of the first domain and to the N-terminal extremity of second domain;

and direct equivalents thereof.

As it is well known, minor changes in an amino acid sequence such as deletion, addition or substitution of one or several amino acids may lead to an allelic form of the original protein which has substantially identical properties. Thus, by the term "direct equivalents thereof" is meant either any single domain CD25 binding molecule (molecule X)

- (i) in which the hypervariable regions CDR1, CDR2 and CDR3 taken as a whole are at least 80% homologous, pref rably at least 90% homologous, more preferably at least 95% homologous to the hypervariable regions as shown in Seq. Id. No. 1 and,
- (ii) which is capable of inhibiting the binding of IL-2 to its receptor substantially to the same extent as a

reference molecule having framework regions identical to those of molecule X but having hypervariable regions CDR1, CDR2 and CDR3 identical to those shown in S q. Id. No. 1;

or any CD25 binding molecule having at least two domains per binding site (molecule X')

- (i) in which the hypervariable regions CDR1, CDR2, CDR3, CDR1', CDR2' and CDR3' taken as a whole are at least 80% homologous, preferably at least 90% homologous, more preferably at least 95% homologous to the hypervariable regions as shown in Seq. Id. No. 1 and 2 and
- (ii) which is capable of inhibiting the binding of IL-2 to its receptor substantially to the same extent as a reference molecule having framework regions and constant parts identical to molecule X' but having hypervariable regions CDR1, CDR2, CDR3, CDR1', CDR2' and CDR3' identical to those shown in Seq. Id. No. 1 and 2.

This last criterion may be conveniently tested in various assays including a Mixed Lymphocyte Reaction (MLR) bioassay, an antigen specific HPBM response bioassay and an IL-2 dependent T lymphoblast proliferation bioassay. Such assays are described hereinafter in the text. By the term "to the same extent" is meant that the reference and the equivalent molecules exhibit, on a statistical basis, essentially identical IL-2 binding inhibition curves in one of the bioassays referred to above.

Most preferably, the chimeric CD25 antibody comprises at least

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a) one heavy chain which comprises a variable domain having an amino acid sequence substantially identical to that shown in Seq. Id. No. 1 starting with amino acid at position 1 and ending with amino acid at position 117 and the constant part of a human heavy chain; and

b) one light chain which comprises a variable domain having an amino acid sequence substantially identical to that shown in Seq. Id. No. 2 starting with glutamic acid at position 1 and ending with glutamic acid at position 104 and the constant part of a human light chain.

The constant part of a human heavy chain may be of the  $\gamma_1$ ,  $\gamma_2$ ,  $\gamma_3$ ,  $\gamma_4$ ,  $\mu$ ,  $\alpha_1$ ,  $\alpha_2$ ,  $\delta$  or  $\epsilon$  type, preferably of the  $\gamma$  type, more preferably of the  $\gamma_1$  type, whereas the constant part of a human light chain may be of the  $\kappa$  or  $\lambda$  type (which includes the  $\lambda_1$ ,  $\lambda_2$  and  $\lambda_3$  subtypes) but is preferably of the  $\kappa$  type. The amino acid sequence of all these constant parts are given in Kabat et al (Supra).

Conjugates of the CD25 binding molecules of the invention, e.g. enzyme or toxin or radioisotope conjugates, are also included within the scope of the invention.

A CD25 binding molecule of the invention may be produced by recombinant DNA techniques. In view of this, one or more DNA molecules encoding the binding molecule must be constructed, placed under appropriate control sequences and transferred into a suitable host organism for expression.

In a very general manner, there are accordingly provided

- (i) DNA molecules encoding a single domain CD25 binding molecule, of the invention, a single chain CD25 binding molecule of the invention, a heavy or light chain or fragments thereof of a CD25 binding molecule of the invention and
- (ii) the use of the DNA molecules of the invention for the production of a CD25 binding molecule of the invention by recombinant means.

The present state of the art is such that the skilled man will be able to synthetize the DNA molecules of the invention given the information provided herein i.e. the amino acid sequences of the hypervariable regions and the DNA sequences coding for them. A method for constructing a variable domain gene is for example described in EPA 239 400 and may be briefly summarized as follows: A gene encoding a variable domain of a MAb of whatever specificity is cloned. The DNA segments encoding the framework and hypervariable regions are determined and the DNA segments encoding the hypervariable regions are removed so that the DNA segments encoding the framework regions are fused together with suitable restriction sites at the junctions. The restriction sites may be generated at the appropriate positions by mutagenesis of the DNA molecule by standard procedures. Double stranded synthetic CDR cassettes are prepared by DNA synthesis according to the sequences given in Seq. Id. No. 1 or 2. These cassettes are provided with sticky ends so that they can be ligated at the junctions of the framework. A protocol for achieving a DNA molecule encoding an immunoglobulin variable domain is shown in Figure 5.

Furthermore, it is not necessary to have access to the mRNA from a producing hybridoma cell line in order to obtain a DNA construct coding for the MAbs of the invention. Thus PCT application W0 90/07861 gives full instructions for the production of a MAb by recombinant DNA techniques given only written information as to the nucleotide sequence of the gene. The method comprises the synthesis of a number of oligonucleotides, their amplification by the PCR method, and their splicing to give the desired DNA sequence.

Expression vectors comprising a suitable promoter or genes encoding heavy and light chain constant parts are publicly available. Thus, once a DNA molecule of the invention is prepar d it may be conveniently transferred in an appropriate expr ssion vector. DNA molecules encoding singl chain antibodies may also b prepared by standard methods, for example, as d scribed in W0 88/1649.

In view of the foregoing, and since the mouse MAb as naturally secreted by the hybridoma is not the preferred type of MAb, it is believed that no hybridoma d posit is necessary to comply with the criteria of sufficiency of description.

In a particular embodiment of the invention, the recombinant means for the production of a CD25 binding molecule includes first and second DNA constructs as described below:

The first DNA construct encodes a heavy chain or fragment thereof and comprises

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a) a first part which encodes a variable domain comprising alternatively framework and hypervariable regions, said hypervariable regions being in sequence CDR1, CDR2 and CDR3 the amino acid sequences of which are shown in Seq. Id. No. 1; this first part starting with a codon encoding the first amino acid of the variable domain and ending with a codon encoding the last amino acid of the variable domain, and b) a second part encoding a heavy chain constant part or fragment thereof which starts with a codon encoding the first amino acid of the constant part of the heavy chain and ends with a codon encoding the last amino acid of the constant part or fragment thereof, followed by a non-sense codon.

Preferably, this first part encodes a variable domain having an amino acid sequence substantially identical to the amino acid sequence as shown in Seq. Id. No. 1 starting with the amino acid at position 1 and ending with the amino acid at position 117. More preferably the first part has the nucleotide sequence as shown in Seq. Id. No. 1 starting with the nucleotide at position 142 and ending with the nucleotide at position 492. Also preferably, the second part encodes the constant part of a human heavy chain, more preferably the constant part of the human  $\gamma$ 1 chain. This second part may be a DNA fragment of genomic origin (comprising introns) or a cDNA fragment (without introns).

The second DNA construct encodes a light chain or fragment thereof and comprises

a) a first part which encodes a variable domain comprising alternatively framework and hypervariable regions; said hypervariable regions being in sequence CDR1', CDR2' and CDR3', the amino acid sequences of which are shown in Seq. Id. No. 2; this first part starting with a codon encoding the first amino acid of the variable domain and ending with a codon encoding the last amino acid of the variable domain, and b) a second part encoding a light chain constant part or fragment thereof which starts with a codon encoding the first amino acid of the constant part of the light chain and ends with a codon encoding the last amino acid of the constant part or fragment thereof followed by a non-sense codon.

Preferably, this first part encodes a variable domain having an amino acid sequence substantially identical to the amino acid sequence as shown in Seq. Id. No. 2 starting with the amino acid at position 1 and ending with the amino acid at position 104. More preferably, the first part has the nucleotide sequence as shown in Seq. Id. No. 2 starting with the nucleotide at position 244 and ending with the nucleotide at position 555 Also preferably the second part encodes the constant part of a human light chain, more preferably the constant part of the human  $\kappa$  chain.

In the first and second DNA constructs, the first and second parts are preferably separated by an intron. In the intron located between the first and second part, an enhancer is preferably inserted. The presence of this genetic element which is transcribed but not translated, may be required for an efficient transcription of the second part. More preferably the first and second DNA constructs comprise the enhancer of a heavy chain gene advantageously of human origin.

The first or second DNA construct advantageously comprises a third part which is located upstream of the first part and which encodes part of a leader peptide; this third part starting with the codon encoding the first amino acid and ending with the last amino acid of the leader peptide. This peptide is required for secretion of the chains by the host organism in which they are expressed and is subsequently removed by the host organism. Preferably, the third part of the first DNA construct encodes a leader peptide having an amino acid sequence substantially identical to the amino acid sequence as shown in Seq. Id. No. 1, starting with the amino acid at position -19 and ending with the amino acid at position -1. Also preferably, the third part of the second DNA construct encodes a leader peptide having an amino acid sequence as shown in Seq. Id. No. 2, starting with the amino acid at position -22 and ending with the amino acid at position -1.

Each of the DNA constructs are placed under the control of suitable control sequences, in particular under the control of a suitable promoter. Any kind of promoter may be used, provided that it is adapted to the host organism in which the DNA constructs will be transferred for expression. However, if expression is to take place in a mammalian cell, it is particularly preferred to use the promoter of an immunoglobulin gene.

The desired antibody may be produced in a cell culture or in a transgenic animal. A suitable transgenic animal may be obtained according to standard methods which include micro injecting into eggs the first and second DNA constructs placed under suitable control sequences transferring the so prepared eggs into appropriat pseudo-pregnant females and selecting a descendant expressing the desired antibody.

When the antibody chains have to be produced in a cell culture, the DNA constructs must first be inserted into either a single expression vector or into two separate but compatible expression vectors, the latter possi-

bility being preferred.

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Accordingly, the invention also provides an expression vector able to replicate in a prokaryotic or ukaryotic cell line which comprises at least one of the DNA constructs abov described.

Each expression vector containing a DNA construct is then transferred into a suitable host organism. When the DNA constructs are separately inserted on two expression vectors, they may be transferred separately, i.e. one type of vector per cell, or co-transferred, this latter possibility being preferred. A suitable host organism may be a bacterium, a yeast or a mammalian cell line, this latter being preferred. More preferably, the mammalian cell line is of lymphoid origin e.g. a myeloma, hybridoma or a normal immortalized B-cell, but does not express any endogeneous antibody heavy or light chain.

It is also preferred that the host organism contains a large number of copies of the vectors per cell. If the host organism is a mammalian cell line, this desirable goal may be reached by amplifying the number of copies according to standard methods. Amplification methods usually consist of selecting for increased resistance to a drug, said resistance being encoded by the expression vector.

In another aspect of the invention, there is provided a process for producing a multi-chain CD25 binding molecule which comprises (i) culturing an organism which is transformed with first and second DNA constructs of the invention and (ii) recovering an active CD25 binding molecule from the culture.

Alternatively, the heavy and light chains may be separately recovered and reconstituted into an active binding molecule after in vitro refolding. Reconstitution methods are well-known in the art; Examples of methods are in particular provided in EPA 120 674 or in EPA 125 023.

Therefore a process may also comprise

- (i) culturing a first organism which is transformed with a first DNA construct of the invention and recovering said heavy chain or fragment thereof from the culture and
- (ii) culturing a second organism which is transformed with a second DNA construct of the invention and recovering said light chain or fragment thereof from the culture and
- (iii) reconstituting in vitro an active CD25 binding molecule from the heavy chain or fragment thereof obtained in (i) and the light chain or fragment thereof obtained in (ii).

In a similar manner, there is also provided a process for producing a single chain or single domain CD25 binding molecule which comprises (i) culturing an organism which is transformed with a DNA construct respectively encoding a single chain or single domain CD25 binding molecule of the invention and (ii) recovering said molecule from the culture.

CD25 binding molecules of the invention exhibit very good immunomodulatory activity as shown, for example, in a mixed lymphocyte reaction (MLR) bioassay (Akbar et al, J. Immunol. 140, 2171-8). The MLR is generally considered to be the <u>in vitro</u> equivalent of the allogeneic transplant response which leads to rejection <u>in</u> vivo.

## 1. Inhibition of the MLR

From a HPBM preparation of a first donor are aliquoted 100  $\mu$ l samples containing 10<sup>5</sup> HPBM to which are added various concentrations of a molecule of the invention ranging from 0 to 300 ng/ml (including these limiting values). Then each sample is mixed with a 100  $\mu$ l aliquot containing 10<sup>5</sup> HLA-incompatible X-irradiated HPBM of a second donor, or T-cell depleted HPBM. The mixture is incubated for 6 days at 37°C, and 1  $\mu$ Ci of methyl <sup>3</sup>H-thymidine (<sup>3</sup>H-Tdr) in 10  $\mu$ l volume is then added. After 6 hours, the cell proliferation is measured by radio-activity incorporation.

In this particular assay, the molecules of the invention show an <u>in vitro</u> immunomodulatory activity at concentrations of from 0.3 ng/ml as shown in Figure 6. 50% of the cellular growth is inhibited at about 3 ng/ml.

The immunomodulatory activity of the molecules of the invention may also be estimated by measuring the inhibition of antigen-specific HPBL response or the inhibition of IL-2 dependent T-lymphoblast proliferation as follows:

## 2. Inhibition of antigen-specific HPBM response

The molecules of the invention inhibit efficiently the generation of a PPD (tuberculin) specific, HLA class II restricted T-cell response, indicating their ability to inhibit the binding of endogenously produced IL-2 to its receptor. In vivo these antigen specific responses are expected to play a crucial role in the initiation of autoimmunity and transplantation rejection.

From a preparation of HPBM are aliquoted 100 µl samples containing 10<sup>5</sup> HPBM to which are added various concentrations of a molecule of the invention ranging from 0 to 300 ng/ml (including these limiting values) and tuberculin (PPD) at a final concentration of 30 µg/ml. The samples are incubated for 6 days at 37°C, and 1 µCi

of methyl <sup>3</sup>H-thymidine is then added in a 10 µl volume. After 6 hours of incubation, cell proliferation is measured by radioactivity incorporation.

In this particular assay, the molecules of the invention show an immunomodulatory activity of from about 10 ng/ml as shown in Figure 7. 50% of the cellular growth is inhibited at about 50 ng/ml.

#### 3. Inhibition of IL-2 dependent T-lymphoblast proliferation

The molecules of the invention inhibit efficiently the IL-2 dependent growth of human T cell blasts induced by MLR or PPD stimulation. These cells are expected to play a major role in the chronicity of autoimmunity and rejection episodes.

Triplicate cultures containing  $20 \times 10^3$  5 day old PPD or MLR stimulated HPBM in a final volume of  $200 \, \mu$ l are incubated at  $37^{\circ}$ C for 48 hours in the presence of 5, 10 or 20 ng/ml recombinant IL-2 and a molecule of the invention at various concentrations ranging from 0 to 10  $\mu$ g/ml (including these limiting values). Then  $^3$ H-Tdr is added. After 6 hours, cell proliferation is measured by radioactivity incorporation. In this particular assay, the molecules of the invention show an immunomodulatory activity at concentrations of from 0.1  $\mu$ g/ml as shown in Figures 8A and 8B, 8C and 8D.

Therefore the invention also provides

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- (i) the use of a CD25 binding molecule of the invention in immunosuppression of a human immune system (ii) a method of immunosuppressing the human immune system which comprises administering an immunosuppressive effective amount of a CD25 binding molecule of the invention to a patient in need of such treatment.
- (iii) a pharmaceutical composition for immunosuppressing the human immune system which comprises a CD25 binding molecule of the invention and a pharmaceutically acceptable carrier or diluent.

In particular, a CD25 binding molecule of the invention is useful for preventing or treating graft rejection episodes.

For these indications, the appropriate dosage will, of course, vary depending upon, for example, the particular molecule of the invention to be employed, the host, the mode of administration and the nature and severity of the condition being treated. However, in prophylactic use, satisfactory results are generally indicated to be obtained at daily dosages from about 0.1 mg to about 1 mg per kilogram body weight. These dosages should be increased by up to a factor of 4 for treating a rejection event when it actually occurs. A molecule of the invention is conveniently administered parenterally, normally intravenously, for example, into the antecubital or other peripheral vein. A prophylactic treatment typically comprises administering the molecule of the invention once daily to once weekly for 2 to 4 weeks, starting on the day of transplantation, preferably some hours before transplantation.

The molecules of the invention may also be useful in the treatment of malignancies of cells expressing the CD25 antigen, for example in the treatment of T-cell leukemia and certain other leukemias and lymphomas. For this purpose, the CD25 binding molecule may be used in the form of a radioconjugate in which the molecule is coupled to an alpha-emitting radionuclide.

The molecules of the invention may also be useful in the treatment or prophylaxis of HIV infection. It appears that the HIV virus requires proliferating T cells in order to multiply, and thus inhibition of T cell proliferation by blocking the CD25 antigen should also inhibit the multiplication of the virus.

Pharmaceutical compositions of the invention may be manufactured in conventional manner. A composition according to the invention is preferably provided in lyophilized form. For immediate administration it is dissolved in a suitable aqueous carrier, for example sterile water for injection or sterile buffered physiological saline. If it is considered desirable to make up a solution of larger volume for administration by infusion rather as a bolus injection, it is advantageous to incorporate human serum albumin or the patient's own heparinised blood into the saline at the time of formulation. The presence of an excess of such physiologically inert protein prevents loss of monoclonal antibody by adsorption onto the walls of the container and tubing used with the infusion solution. If albumin is used, a suitable concentration is from 0.5 to 4.5% by weight of the saline solution.

According to a further aspect of the invention, it has been found that particularly beneficial results can be achieved by the use, in combination, of at least two antigen binding molecules to activated T-cells, said binding molecules recognizing at least two different antigens characteristic of activated T-cells.

Preferably a combination of two different antigen binding molecules is used, each recognizing a different antigen. Thus although both antigen binding molecules recognize activated T-cell surface antigens, they are not competing with each other for the same binding site on activated T-cells.

Preferably one of th antigen binding molecules is a CD25 binding molecule.

Accordingly the present invention also provides an immuno-suppressive composition comprising a mixture of at I ast one CD25 binding molecul and at least one antigen binding molecule to at least one antigen other

than CD25 which is characteristic of activated T-cells.

The invention further provides at least two antigen binding molecules to activat d T-c IIs in association with one another for use in immunosuppression of the mammalian system, said antigen binding molecules recognizing at least two different antigens characteristic of activated T-cells, one of which is the CD25 antigen.

By "antigen binding molecule to activated T-cells" is meant a binding molecule which strongly reacts with activated T-cells while it reacts weakly or not at all with resting T-cells. Preferably the antigen binding molecules are complete immunoglobulin molecules, more preferably murine, chimeric, or humanized monoclonal antibodies, particularly chimeric monoclonal antibodies. The preferred CD25 monoclonal antibodies are those having CDR's with the amino acid sequences described above.

Advantageously, the composition of the invention may also include or may be used in combination with an immunosuppressive drug such as cyclosporin A.

The preferred monoclonal antibodies to activated T-cell antigens other than CD25 are typically those classified in the CD7 cluster as established by the Boston Workshop and reported in "Leucocyte Typing II, Vol. I human T lymphocytes" by Reinherz, Haynes, Nadler and Berstein, Springer Verlag, 1985. The CD7 antigen is heterogeneously expressed on about 80% of resting T-cells. However, the expression strongly increases upon activation (a 2-3 fold rise in intensity).

A preferred combination of antibodies is therefore a combination of a CD7 with a CD25 antibody. Accordingly, the composition of the invention preferably comprises a mixture of at least one CD25 antibody together with at least one CD7 antibody, more preferably of one CD25 antibody together with one CD7 antibody. Also preferably, both antibodies are of the IgG isotype.

The two antibodies, optionally together with an immunosuppressive drug, can be used in clinical practice in various ways. Preferably they are mixed together and the physical mixture is administered to the patient. An alternative procedure is the administration of the antibodies and optionally the immunosuppressive drug to the recipient from separate reservoirs in any order but at the same time. The composition may be prepared and administered parenterally as described above for the single CD25 antibody. Alternatively, the immunosuppressive drug is administered orally and the monoclonal antibodies are administered parenterally, separately or as a mixture.

To aid in making up suitable compositions, the monoclonal antibodies and optionally an immunosuppressive drug, may be packaged separately within the same container, with instructions for mixing or concomitant administration. Examples of kits include for example a multi-barrelled syringe or a twin pack containing separate unit dose forms of at least two antibodies to activated T-cells, said antibodies recognizing at least two different antigens characteristic of activated T-cells, one of which is the CD25 antigen.

Investigations so far indicate that the administration of the antibodies in combination with one another and optionally with an immunosuppressive drug is free from unacceptable side-effects at the dosage levels employed and that there is no potentiation of the side-effects observed with the individual antibodies. For use in prophylaxis, a suitable dosage will normally call for the administration of the order of 0.05 - 0.5 milligram of a first antibody (such as the CD25 antibody) per kilogram body weight of the patient and 0.05 - 0.5 milligram of a second antibody (such as a CD7 antibody) per kilogram body weight. When the immunosuppressive drug is cyclo-sporin, the recommended amount of the immunosuppressive drug which can be optionally used is 2 to 5 milligram per kilogram body weight when administered parenterally and 10 - 15 mg/kilogram body weight when administered orally. The composition of the invention may be administered on a daily or weekly basis, preferably on a weekly basis.

Although the composition of the invention is particularly designed for use in prophylaxis of graft rejection episodes, its use can be conveniently extended to the treatment of rejection events when they actually occur. In this case, the dosages should be increased by up to a factor of 4.

Murine monoclonal antibodies suitable for use in the present invention are known per se. Many monoclonal antibodies against activated T-cell surface antigens are available from Culture Collections in various countries of the world and specifically, the American Type Culture Collection of Rockville, Maryland, USA can provide suitable monoclonal antibodies or hybridomas secreting such antibodies. An example of hybridoma secreting CD7 monoclonal antibodies that can be used in the present invention and that is available from the ATCC is T3-3A1. Other CD7 antibodies are RFT-2 and CHH 380 (a chimeric antibody). CD25 antibodies include, besides the preferred RFT-5 and its chimeric derivative as described above; M7/2 (Gaulton et al, Clin. Immunol. and Immunopath. (1985) 36: 18); the anti-tac antibody (Uchiyama et al, J. Immunol. (1981) 126 (4): 1393); and the Campath 6 monoclonal antibody.

The synergistic effect of a combination of CD25 and CD7 monoclonal antibodies is demonstrated in vitro by the MLR bioassay described abov , and also in vivo in clinical tests on human patients.

In the MLR bioassay, inhibition of th <sup>3</sup>H-TdR uptake is observed in cultur s to which a CD7 (RFT2) or a CD25 (RFT5) monoclonal antibody are added singly, and ther is a substantially gr ater d gree of inhibition

when both of these antibodies are used together at the same total concentration. The MLR is the <u>in vitro</u> equivalent of the allogeneic transplant response which leads to rejection <u>in vivo</u> while the inhibition described above is equivalent to immunosuppression <u>in vivo</u>.

In MLRs to which cyclosporin is added over a dose range from 10 nanograms/ml to 100 µg/ml, in the presence of CD7 or CD25 monoclonal antibodies there is an increased inhibition of <sup>3</sup>H-TdR compared to cyclosporin alone over the whole dose range. The combination of CD7, CD25 and cyclosporin shows a greater inhibitory effect than any other combination.

In clinical tests, patients about to undergo kidney, liver or heart transplantation are selected for prophylactic therapy. On the day of transplantation, 2 hours before surgery, a first intravenous infusion of the chimeric CD25 antibody of Example 5 together with chimeric CD7 antibody (CHH 380) is administered at a dose of 0.2 mg of each antibody per kg of body weight. Two days after surgery an identical infusion of the two antibodies at 0.4 mg/kg of body weight is administered and then repeated at weekly intervals for one month.

The intravenous infusions are prepared as follows: the lyophylized antibodies are mixed together and dispersed into 100 ml sterile buffered saline containing 4.5% wt. of human albumin. This saline dispersion is administered to the patients over a 30 minute period. The patients also receive standard cyclosporin therapy. No patients undergo a rejection episode during the one month therapy period.

#### Brief description of the drawings

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Figure 1A is a schematic diagram showing the structure of an IgG molecule as well as the genes encoding the heavy and light chains. Figure 1B schematically represents the arrangement of a variable domain of a heavy or light chain into framework (FR) and hypervariable (CDR) regions.

Figures 2A and 2B show the analysis of EcoRI-digested genomic DNA of mouse hybridoma RFT5-IgG2a (1), RFT5-IgG1 (2), RFT4 (3) and NS-1 (4) by Southern blot using a  $^{32}$ P labelled DNA probe either encoding the murine heavy chain enhancer (Fig. 2A) or encoding the mouse  $C_k$  and the five  $J_k$  gene segments (Fig. 2B). 10  $\mu$ g of genomic DNA are digested with EcoRI and are size-fractionated on a 0.8% agarose gel. Then the fragments are transferred on a nitrocellulose membrane and hybridized with the probe. After washing, the membrane is exposed overnight on a Kodak X-0 Mat film.

Figures 3A and 3B show the parental expression vectors pSV2-neo-huC $\gamma$ 1 and pSV2-DHFR-E $\mu$ -huC $\kappa$ . Both plasmids comprise an ampicillin resistance gene (ampR) and the origin of replication of pBR322 and SV40 (pBR322 ori and SV40 ori). pSV2-neo-huC $\gamma$ 1 is characterized by the presence of a neomycin gene (neoR) and the gene encoding the human  $\gamma_1$  constant part (huC $\gamma_1$ ), while pSV2-DHFR-E $\mu$ -huC $\kappa$  has inserted a dihydro-folate reductase (DHFR) gene (methotrexate resistance) and the gene encoding the human  $\kappa$  constant part (huC $\kappa$ ). The final vectors for expressing the chimeric heavy or light chain are respectively obtained by inserting into pSV2-neo-hC $\gamma$ 1 a DNA fragment encoding the leader peptide (L), and the variable domain (VDJ<sub>2</sub>) of the RFT5-lgG2a heavy chain together with the human heavy chain enhancer and by inserting into pSV2-DHFR-E $\mu$ -huC $\kappa$  a DNA fragment encoding the leader peptide (L) and the variable domain (VJ<sub>2</sub>) of the RFT5-lgG2a light chain.

Figures 4A and 4B show the productivity of individual cell pools grown at increasing concentration of methotrexate (MTX) respectively according to procedures A and B described in Example 5. The Y-axis of the graph gives the amount of monoclonal antibody produced in mg/10° cells in 72 hours.

Figure 5 shows a protocol for constructing CDR replacements by insertion of CDR cassettes into a vector containing 4 framework regions fused together.

Figure 6 shows the inhibition of MLR by (x) RFT5-IgG2a ( $\gamma_2 a$ ,  $\kappa$ ) and (o) a murine-human chimeric MAb of the invention ( $\gamma_1$ ,  $\kappa$ ). Both MAbs have the variable domains as shown in Seq. Id. No. 1 and 2.

Figure 7 shows the inhibition of PPD specific HPBM response by (x) RFT5-IgG2a and (o) the same murine-human chimeric MAb.

Figure 8 shows the effect of RFT5-IgG2a and of the same murine-human chimeric MAb on PPD T-lymphoblast proliferation, (Fig. 8B and 8A) and on MLR T-lymphoblast proliferation, (Fig. 8D and 8C) at an IL-2 concentration of 5 ng/ml (o), 10 ng/ml ( ) and 20 ng/ml (x).

By way of illustration only, the production of a chimeric CD25 antibody of the invention is exemplified as follows:

## Example 1 Cloning of the gene encoding the variable domain of the heavy chain of RFT5-IgG2a

The genomic DNA of the hybridomas RFT5-IgG2a (CD25;  $\gamma_{2a}$ ;  $\kappa$ ), RFT5-IgG1 (CD25;  $\gamma_1$ ;  $\kappa$ ) and RFT4 (CD4;  $\gamma_1$ ,  $\kappa$ ) and of the parental myeloma c  $\,$  Il line of the hybridomas, namely NS-1, is isolated and digest  $\,$  d with EcoRl. Each digested DNA is then fractionated on the same agarose gel. After migration, the agaros  $\,$  gel is analysed

by Southern blot using as probe a <sup>32</sup>P-labelled 0.7kb Xbal-EcoRI DNA fragment which encodes the murine heavy chain enhancer (Heinrich et al, J. of Immunol. (1989) 143: 3589). 3 types of bands are revealed on the gel after hybridization as shown in Figure 2. The 6.5 kb EcoRI fragment is present in the DNA digest of all cell lines including NS-1, the parental myeloma cell-line and therefore is of no interest. The 2.9 kb EcoRI fragment is only detected in the DNA digest of the hybridoma RFT5-IgG1 and is thought to be the result of an abnormal gene rearrangement. The 6.8 kb EcoRI fragment which is absent in the DNA digest of the parental cell line NS-1 is therefore a fragment of choice and further punification of this fragment is consequently carried out by preparative agarose gel electrophoresis.

DNA fragments of approximately 5-7 kb are cloned in the EcoRI restriction site of bacteriophage ZAP (Stratagene). Using the probe described above, 6x10<sup>6</sup> recombinant phages are screened and 11 clones are found to hybridize. The DNA insert of the 11 clones was amplified on phage plate lysate by polymerase chain reaction (PCR) using as primers, a first oligonucleotide encoding the murine J<sub>2</sub> gene and a second oligonucleotide encoding the beginning of the RFT5 heavy chain up to amino acid No. 7 (sequence previously determined). The second primer is designed taking into account the most frequent codon usage genes. The DNA fragments obtained from each of the 11 clones are analysed by Southern blot using as probe an oligonucleotide encoding the amino acid sequence comprised between amino acids 20 and 27 of the RFT5 heavy chain which is also designed according to the most frequent codon usage. 9 identical phage clones are revealed using the probe. Part of the DNA insert which encodes the variable domain is sequenced by the dideoxy termination method and is to be seen in Seq. Id. No. 1.

## Example 2 Construction of a chimeric RFT5 heavy chain gene

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A 6kb EcoRI fragment obtained by digestion of the DNA of one of the 9 phage clones and comprising the gene of the RFT5 heavy chain variable domain (including the promoter and the enhancer) is cloned into the EcoRI restriction site of the eukaryotic expression vector pSV2 neo-human  $\gamma_1$  constant part (Heinrich et al, supra) as shown in Figure 3A. Then the nucleotide sequence of the gene encoding the RFT5 heavy chain variable domain is redetermined to exclude the possibility that a mutation in this gene has occurred during the propagation of the plasmid.

## Example 3 Cloning of the gene encoding the variable domain of the light chain of RFT5

The genomic DNA of the hybridomas RFT5, RFT5\* and RFT4 and of the parental cell line NS-1 is isolated and digested with EcoRI. Each digested DNA is then fractionated on the same agarose gel. After migration, the agarose gel is analysed by Southern blot using as probe a  $^{32}P$  labelled DNA fragment comprising the 5 mouse  $J_{\kappa}$  genes and the mouse  $C_{\kappa}$  gene.

3 major types of bands of approximately 12, 16 and 18 kb are revealed on the gel after hybridization, as shown in Figure 2B. The largest fragments are the only ones specific for the RFT5 hybridoma. Size fractionated EcoRl fragments of approximately 18 kb are cloned in phage EMBL4 (Stratagene) 7x10<sup>5</sup> recombinant phage clones are screened with the probe described above and 2 clones are found to hybridize each comprising an identical 18 kb insert. A 4.4 kb EcoRl - Xbal subfragment is shown to contain the full gene encoding the RFT5 light chain variable domain and is cloned into the plasmid pGEM4 (Stratagene). The sequence of the 4.4 kb fragment is determined. Part of the 4.4 kb DNA insert which encodes the variable domain is sequenced. The sequence is to be seen in Seq. Id. No. 2.

## Example 4 Construction of a chimeric RFT5 light chain gene

A 1.1 kb Xbal - Xbal fragment encoding the murine heavy chain enhancer (Heinrich et al; supra) together with a HindIII - SphI fragment encoding the human  $\kappa$  constant part is subcloned in phage mp18 (Stratagene). After disruption of restriction sites by mutagenesis a filled-in EcoRI - HindIII fragment comprising the sequence for the murine heavy chain enhancer (E $\mu$ ) and the human  $\kappa$  constant part (huC $\kappa$ ) is cloned in the filled in EcoRI - BamHI site of pSV2-DHFR. pSV2-DHFR is obtained by replacing the BamHI - HindIII fragment of pSV2-neo with a BamHI - HindIII fragment encoding the DHFR gene.

The 4.4 kb EcoRI - Xbal fragment of Example 3 is then inserted into pSV2-DHFR-Εμ-huCκ.

## Example 5 Expression of a RFT5 chimeric antibody

The plasmids as obtained in Examples 2 and 4 are co-transferred in the mouse myeloma cell line SP2/0 (ATCC CRL 1581) by electroporation using a gene puls r apparatus from Biorad. This technique is known to

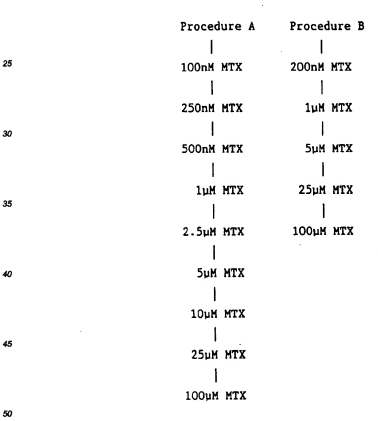
create stable transfectants at a high frequency. The SP2/0 cell line fails to produce endogeneous heavy and light chains and is sensitive to Geneticin (G 418) at a concentration of 0.8 mg/l.

SP2/0 cells are grown in the usual growth medium (RPMI + 10% FCS +  $5x10^{-6}$   $\beta$ -mercaptoethanol) harvested in the log phase of growth and washed with the electroporation buffer (Bio-Rad). Cell concentration is adjusted to  $2x10^7$  cells/ml. To 0.8 ml of the cell suspension is added 15-20  $\mu$ g of each plasmid. The mixture is placed on ice and left to stand for 10 min. Then the cells are subjected to an electrical pulse (280 Volt; 25  $\mu$ F) and again left to stand for 15 min. Cells are transferred to the usual growth medium and incubated at 37°C in a CO<sub>2</sub> incubator.

After 3-day incubation, selection for G 418 resistance is started. Cells are resuspended in fresh medium containing 1.4 mg/ml G 418. The cultures yield growing cells after 10-14 day-incubation in the presence of G 418. After 2-week incubation, the supernatants of the confluent cultures are tested for human  $\lg G$  expression in a sandwich-type ELISA (anti-human  $\kappa$ -light chain / supernatant /anti-human  $\lg G$ -alkaline phosphatase conjugate).

This test indicates that complete antibody molecules are secreted in all cultures at varying concentrations in the range of 50-500 ng/ml.

To select cells in which the DHFR gene is amplified and therefore secrete high amounts of the desired antibody two selection procedures for Methotrexate (MTX) resistance are carried out as described below. For this purpose, the G 418 resistant cell pools are each divided and amplification is proceeded either according to procedure A (MTX increase by a factor of 2 or 2.5) or procedure B (MTX increase by a factor of 5).



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Each amplification step comprises inoculating the cells at a density of 2x10<sup>5</sup> cells/ml in the usual growth medium supplemented with G 418 at 1.4 mg/ml and with MTX at the concentration of choice. After 72 hour incubation, cells and the supernatant are separated. Antibody secretion is monitored either by ELISA or by HPLC using a protein A column.

Figures 4A and 4B show the antibody productivity of some transfectant pools. Most of the pools reach a maximum of specific antibody production at a certain MTX concentration. The best producing pools are cloned by limiting dilution. Out of several hundred analysed clones, 15 best producing clones are selected. Productivity of the clones ranges from 30 to 50 mg MAb/10<sup>9</sup> cells in 72 hours.

The antibody is purified from a culture supernatant by elution on a protein A affinity column.

5		SEQUENCE IDENTIFIER No.1
	Subject matter:	The immunoglobulin heavy chain variable domain of the RFT5 antibody
10	Sequence type:	Nucleotide sequence and its corresponding amino acid sequence
	Molecule type:	Genomic DNA
15	Length:	492 nucleotides
	Original source:	A murine hybridoma
	Features of the	nucleotide sequence:
20	An intron i	s located from nucleotide 47 to 130
	D segment g	gene: from nucleotide 142 to 435 gene: from nucleotide 436 to 447 gene: from nucleotide 448 to 492
25	Features of the	amino acid sequence:
	Leader pept	ride : from amino acid (a.a.) -19 to -1
30	CDR1: from FR2: from	a.a. 1 to 30 a.a. 31 to 35 a.a. 36 to 49 a.a. 50 to 66
35	FR3: from CDR3: from	a.a. 67 to 98 a.a. 99 to 106 a.a. 107 to 117.
40	ATG GAA TGT AAC	TGG ATA CTT CCT TTT ATT CTG TCG GTA ATT TCA G 46
	Met Glu Cys Asn	Trp Ile Leu Pro Phe Ile Leu Ser Val Ile Ser -15 -10 -5
		-13 -10 -3
45	GTAAGGGGCT CACCA	AGTTCC ATATCTGAAA GAGGATACAG GGTCTGAAGT GACAATGACA 106
	TCTACTCTGC TGTT	CTCTCC ACAG GG GTC TAC TCA GAG GTT CAG CTC CAG 156
50		Gly Val Tyr Ser Glu Val Gln Leu Gln

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-1 1

	CAG	TCT	GGG	ACT	GTG	CTG	GCT	AGG	CCT	GGG	GCT	TCC	GTG	AAG	ATG	TCC	204
5	Gln	Ser	Glu	Thr	Val	Leu	Ala	Arg	Pro	Gly	Ala	Ser	Val	Lys	Met	Ser	
					10					15					20		
10	TGC	AAG	GCT	TCT	GGC	TAC	AGC	TTT	ACC	AGG	TAC	TGG	ATG	CAC	TGG	ATA	252
	Cys	Lys	Ala	Ser	Gly	Tyr	Ser	Phe	Thr	Arg	Tyr	Trp	Met	His	Trp	Ile	
				25					30					35			
15	AAA	CAG	AGG	CCT	GGA	CAG	GGT	CTA	GAA	TGG	ATT	GGT	GCT	ATT	TAT	CCT	300
	Lys	Gln	Arg	Pro	Gly	Gln	Gly	Leu	Glu	Trp	Ile	Gly	Ala	Ile	Tyr	Pro	
			40					45					50				
20	GGA	AAT	ACT	GAT	ACT	AGT	TAC	AAC	CAG	AAG	TTC	GAG	GGC	AAG	GCC	AAA	348
	Gly	Asn	Ser	Asp	Thr	Ser	Tyr	Asn	Gln	Lys	Phe	Glu	Gly	Lys	Ala	Lys	
		55					60					65					
25																	
	ÇTG	ACT	GCA	GTC	ACA	TCC	GCC	AGC	ACT	GCC	TAC	ATG	GAG	CTC	AGC	AGC	396
	Leu	Thr	Ala	Val	Thr	Ser	Ala	Ser	Thr	Ala	Tyr	Met	Glu	Leu	Ser	Ser	
	70			,		75					80					85	
30																	
	CTG	ACA	CAT	GAG	GAC	TCT	GCG	GTC	TAT	TAC	TGT	TCA	AGA	GAC	TAC	GGC	444
	Leu	Thr	His	Glu	Asp	Ser	Ala	Val	Tyr	Tyr	Cys	Ser	Arg	Asp	Tyr	Gly	
35					90	)				95	,				100		
	TAC	TAC	TTT	GAC	TTC	TG0	GGC	CAA	GGC	ACC	ACT	CTC	ACA	GTC	TCC	TCA	492
	Tyr	Туг	Phe	. Asp	Phe	Tr	Gly	Gln	Gly	Thr	The	Leu	Thr	Va]	. Ser	Ser	
40				105	5				110	)				115	j		

5		SEQUENCE IDENTIFIER No. 2												
	Subject matter:	The immunoglobulin light chain variable domain of the antibody	RFT5											
10	Sequence type:	Nucleotide sequence and its corresponding amino acid sequence												
	Molecule type:	Genomic DNA												
15	Length:	455 nucleotides												
	Original source: A murine hybridoma													
	Peatures of the nucleotide sequence:													
20	An intron i	s located from nucleotide 50 to 226												
		ene: from nucleotide 244 to 519 gene: from nucleotide 520 to 455												
25	Features of the	amino acid sequence:												
	leader pept	ide: from a.a22 to -1												
30	CDR1': fro FR2': fro CDR2': fro FR3': fro CDR3': fro	om a.a. 1 to 23 om a.a. 24 to 33 om a.a. 34 to 48 om a.a. 49 to 55 om a.a. 56 to 87 om a.a. 88 to 94 om a.a. 95 to 104.												
35														
		GTG CAG ATT TTC AGC TTC CTG CTA ATC AGT GCC TCA G 49  Val Gln Ile Phe Ser Phe Leu Leu Ile Ser Ala Ser  -15  -10												
40														
	GTAACAGAGG GCAGG	GGAATT TGAGATCAGA ATCCAACCAA AATTATTTTC CCTGGGGAAT 109												
45	TTGAGTCTAA AATAC	CAGTTT TTTTTCTTT TTCTTCATCT GAATGTTGGG TGGTATAAAA 169												
	TTATTTTTGT TTCTC	CTATTT CTACTAATCC CTTTCTCTCT ATTTTGCTTT TTTCTAG 226												
50	TC ATA CTG TCC	AGA GGA CAA ATT GTT CTC ACC CAG TCT CCA GCA ATC 273												
		Arg Gly Gln Ile Val Leu Thr Gln Ser Pro Ala Ile												
	-5	-1 1 5 10												

5	ATG	TCT	GCA	TCT	CCA	GGG	GAG	AAG	GTC	ACC	ATG	ACC	TGC	AGT	GCC	AGC	321
	Met	Ser	Ala	Ser	Pro	Gly	Glu	Lys	Val	Thr	Met	Thr	Cys	Ser	Ala	Ser	
					15					20					25		
10									TAC								369
	Ser	Ser	Ile	Ser	Tyr	Met	Gln	Trp	Tyr	Gln	Gln	Lys	Pro	Gly	Thr	Ser	
				30					35					40			
15																	
									TCC								417
	Pro	Lys	Arg	Trp	Ile	Tyr	Asp	Thr	Ser	Lys	Leu	Ala	Ser	Gly	Val	Pro	
			45					50					55				
20																	
	GCT	CGC	TTC	AGT	GGC	AGT	GGG	TCT	GGG	ACC	TCT	TAT	TCT	CTC	ACA	ATC	465
	Ala	Arg	Phe	Ser	Gly	Ser	Gly	Ser	Gly	Thr	Ser	Tyr	Ser	Leu	Thr	Ile	
25		60	,				65					70					
	AGC	AGC	ATG	GAG	GCT	GAA	GAT	GCT	GCC	ACT	TAT	TAC	TGC	CAT	CAG	CGG	513
	Ser	Ser	Met	Glu	Ala	Glu	Asp	Ala	Ala	Thr	Tyr	Tyr	Cys	His	Gln	Arg	
30	75					80					85					90	
	AGT	AGT	TAC	ACG	TTC	GGA	GGG	GGG	ACC	AAA	CTG	GAA	ATA	AAA	•		555
35	Ser	Ser	Tyr	Thr	Phe	Gly	Gly	Gly	Thr	Lys	Leu	Glu	Ile	Lys			
					95					100	)						

5			n at 10						
10		hain	amino acid 1 to 23 (vith an occasional residue at 0 and a deletion at in $\lambda$ chains)	th possible 7A, B, C,				th possible 95A, B, C,	a possible 06A)
15		Location on the light chain	1 to 23 (vit residue at 0 )	amino acid 24 to 34 (vith possible insertion numbered as 27A, B, C, D, E and P)	35 to 49	50 to 56	57 to 88	amino acid 89 to 97 (vith possible insertions numbered as 95A, B, C, D, E and P)	amino acid 98-107 (vith a possible insertion numbered as 106A)
20		Location on	amino acid 1 occasional r in λ chains)	amino acid insertion n D, E and F)	amino acid 35 to 49	amino acid 50 to 56	amino acid 57 to 88	amino acid insertions D, E and F)	amino acid insertion n
25	Table I						le )	c,	
30		chain	h an 0)	ith mbered		ith mbered	ith possib A, B and C	with possil 100A, B, ( nd K)	
35		the heavy	to 30 (wit) esidue at (	1 to 35 (w ertions nu	6 to 49	O to 65 (w ertions nu d C)	6 to 94 (w umbered 82	5 to 102 ( umbered as H, I, J, a	03 to 113
40		Location on the heavy chain	aminoacid 1 to 30 (with an occasional residue at 0)	amino acid 31 to 35 (with possible insertions numbered as 35A, 35B)	amino acid 36 to 49	amino acid 50 to 65 (with possible insertions numbered as 52A, B and C)	amino acid 66 to 94 (with possible insertions numbered 82A, B and C)	amino acid 95 to 102 (with possible insertions numbered as 100A, B, C, D, E, F, G, H, I, J, and K)	amino acid 103 to 113
45		1	<b>~</b> •	<b>დ ⊆.</b> დ	æ	w <u>-</u> 2 w	æ	а <del>-</del> п	ro.
50		Region	FR1	CDR1	FR2	CDR2	FR3	CDR3	PR4

## 55 Claims

Claims for the following Contra ting States : AT, BE, CH, DE, DK, FR, GB, IT, LU, NL, SE

1. A CD25 binding molecul which comprises at least one antigen binding sit comprising at least one domain

which comprises in sequence, the hypervariable regions CDR1, CDR2 and CDR3; said CDR1 having the amino acid sequence Arg-Tyr-Trp-Met-His, said CDR2 having the amino acid sequence Ala-Ile-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly, and said CDR3 having the amino acid sequence Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe; or direct equivalents thereof.

- A CD25 binding molecule according to claim 1 which comprises at least one antigen binding site comprising:
  - a) a first domain comprising in sequence the hypervariable regions CDR1, CDR2 and CDR3; said CDR1 having the amino acid sequence Arg-Tyr-Trp-Met-His, said CDR2 having the amino acid sequence Ala-Ile-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly, and said CDR3 having the amino acid sequence Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe and,
  - b) a second domain comprising in sequence the hypervariable regions CDR1', CDR2' and CDR3', said CDR1' having the amino acid sequence Ser-Ala-Ser-Ser-Ser-Ile-Ser-Tyr-Met-Gln, said CDR2 having the amino acid sequence Asp-Thr-Ser-Lys-Leu-Ala-Ser, and said CDR3 having the amino acid sequence His-Gln-Arg-Ser-Ser-Tyr-Thr;
  - or direct equivalents thereof.

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- 3. A CD25 binding molecule according to claim 1 which comprises at least one antigen binding site either comprising a domain having the amino acid sequence shown in Seq. Id. No. 1 starting with amino acid at position 1 and ending with amino acid at position 117 or comprising a first domain as described above and a second domain having the amino acid sequence shown in Seq. Id. No. 2, starting with amino acid at position 1 and ending with amino acid at position 104; or direct equivalents thereof.
- 4. A CD25 binding molecule according to claim 2 or 3 which comprises at least
  - a) one immunoglobulin heavy chain or fragment thereof which comprises (i) a variable domain comprising in sequence the hypervariable regions CDR1, CDR2 and CDR3 and (ii) the constant part or fragment thereof of a human heavy chain; said CDR1 having the amino acid sequence Arg-Tyr-Trp-Met-His, said CDR2 having the amino acid sequence Ala-lle-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly, and said CDR3 having the amino acid sequence Asp-Tyr-Gly-Tyr-Phe-Asp-Phe and
  - b) one immunoglobulin light chain or fragment thereof which comprises (i) a variable domain comprising in sequence the hypervariable regions CDR1', CDR2' and CDR3' and (ii) the constant part or fragment thereof of a human light chain; said CDR1' having the amino acid sequence Ser-Ala-Ser-Ser-Ile-Ser-Tyr-Met-Gln, said CDR2' having the amino acid sequence Asp-Thr-Ser-Lys-Leu-Ala-Ser, and said CDR3' having the amino acid sequence His-Gln-Arg-Ser-Ser-Tyr-Thr;
  - or direct equivalents thereof.
- 5. A CD25 binding molecule according to claim 4 which comprises at least
  - a) one heavy chain which comprises a variable domain having the amino acid sequence shown in Seq. Id. No. 1 starting with amino acid at position 1 and ending with amino acid at position 117 and the constant part of a human heavy chain; and
  - b) one light chain which comprises a variable domain having the amino acid sequence shown in Seq.
     Id. No. 2 starting with glutamic acid at position 1 and ending with glutamic acid at position 104 and the constant part of a human light chain;
- or direct equivalents thereof.
  - 6. A CD25 binding molecule according to claim 4 or 5 in which the constant part or fragment thereof of the human heavy chain is of the  $\gamma_1$  type and the constant part or fragment thereof of the human light chain is of the  $\kappa$  type.
  - A DNA molecule which comprises nucleotide sequences encoding in serial order the hypervariable regions CDR1, CDR2 and CDR3 as shown in Seq. Id. n° 1.
- 8. A DNA construct according to claim 7 which encodes a heavy chain or fragment thereof and comprises a) a first part which encodes a variable domain comprising alternatively framework and hypervariable regions, said hypervariable regions being in sequence CDR1, CDR2 and CDR3 the amino acid sequences of which are shown in Seq. Id. No. 1; this first part starting with a codon encoding the first amino acid of the variable domain and ending with a codon encoding the last amino acid of the variable domain, and

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- b) a second part which encodes a heavy chain constant part or fragment thereof which starts with a codon encoding the first amino acid of the constant part of the heavy chain and ends with a codon encoding the last amino acid of the constant part or fragment ther of, followed by a non-sense codon.
- A DNA construct according to claim 8 which comprises a first part which encodes a variable domain having
  the amino acid sequence shown in Seq. Id. No. 1 starting with the amino acid at position 1 and ending
  with the amino acid at position 117; or direct equivalents thereof.
- 10. A DNA construct according to claim 7 which encodes a light chain or fragment thereof and comprises a) a first part which encodes a variable domain comprising alternatively framework and hypervariable regions; said hypervariable regions being in sequence CDR1', CDR2' and CDR3', the amino acid sequences of which are shown in Seq. Id. No. 2; this first part starting with a codon encoding the first amino acid of the variable domain and ending with a codon encoding the last amino acid of the variable domain, and
  - b) a second part which encodes a light chain constant part or fragment thereof which starts with a codon encoding the first amino acid of the constant part of the light chain and ends with a codon encoding the last amino acid of the constant part or fragment thereof followed by a non-sense codon.
- 20 11. A DNA construct according to claim 10 which comprises a first part which encodes a variable domain having the amino acid sequence shown in Seq. Id. No. 2 starting with the amino acid at position 1 and ending with the amino acid at position 104; or direct equivalents thereof.
  - 12. A process for producing a multi-chain CD25 binding molecule which comprises (i) culturing an organism which is transformed with a DNA construct according to claim 8 or 9 and with a DNA construct according to claim 10 or 11 and (ii) recovering an active CD25 binding molecule from the culture.
  - 13. A pharmaceutical composition for immunosuppressing the human immune system, or for treatment of malignancies of CD25<sup>+</sup> cells, or for treatment of HIV infection which comprises a CD25 binding molecule of any of claims 1 to 6 and a pharmaceutically acceptable carrier or diluent.
  - 14. An immunosuppressive composition comprising a mixture of at least one CD25 binding molecule of any of claims 1 to 6 and at least one antigen binding molecule to at least one antigen other than CD25 which is characteristic of activated T-cells.
- 35 15. A composition according to claim 14 which further comprises cyclosporin A.
  - A composition according to claim 14 or claim 15 in which the antigen binding molecule other than the CD25 binding molecule is a CD7 antibody.
  - 17. A composition according to any one of claims 14 to 16 in which the antibodies are chimeric antibodies.
    - 18. A twin pack containing separate unit dose forms of at least two antibodies to activated T-cells, said antibodies recognizing at least two different antigens characteristic of activated T-cells, one of the antibodies being a CD25 binding molecule of any of claims 1 to 6, together with instructions for mixing or concomitant administration.
    - A twin pack according to claim 18 which further comprises a unit dose form of an immunosuppressive drug.
- 20. The use of a CD25 binding molecule of any of claims 1 to 6 for the preparation of a medicament for immunosuppression of the mammalian system in conjunction with an antigen binding molecule recognizing an antigen other than CD25 characteristic of activated T-cells.

## Claims for the following Contracting States : ES, GR

1. A process for the production of a CD25 binding molecule which comprises at least one antigen binding sit comprising at least one domain which comprises in sequence, the hyp rvariable regions CDR1, CDR2 and CDR3; said CDR1 having the amino acid sequence Arg-Tyr-Trp-Met-His, said CDR2 having the amino acid sequence Ala-Ile-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly, and said CDR3 having the amino acid sequence Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe; or direct equivalents thereof, said

process comprising the step of culturing a hybridoma cell line or an organism transformed with a DNA construct comprising nucleotide sequences encoding in serial order said hypervariable regions CDR1, CDR2 and CDR3, and isolating a CD25 binding molecule from the culture.

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- A process according to claim 1 in which the CD25 binding molecule comprises at least one antigen binding site comprising:
  - a) a first domain comprising in sequence the hypervariable regions CDR1, CDR2 and CDR3; said CDR1 having the amino acid sequence Arg-Tyr-Trp-Met-His, said CDR2 having the amino acid sequence Ala-Ile-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly, and said CDR3 having the amino acid sequence Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe and,
  - b) a second domain comprising in sequence the hypervariable regions CDR1', CDR2' and CDR3', said CDR1' having the amino acid sequence Ser-Ala-Ser-Ser-Ser-Ile-Ser-Tyr-Met-Gln, said CDR2' having the amino acid sequence Asp-Thr-Ser-Lys-Leu-Ala-Ser, and said CDR3' having the amino acid sequence His-Gln-Arg-Ser-Ser-Tyr-Thr;
  - or direct equivalents thereof.
- 3. A process according to claim 1 in which the CD25 binding molecule comprises at least one antigen binding site either comprising a domain having the amino acid sequence shown in Seq. Id. No. 1 starting with amino acid at position 1 and ending with amino acid at position 117 or comprising a first domain as described above and a second domain having the amino acid sequence shown in Seq. Id. No. 2, starting with amino acid at position 1 and ending with amino acid at position 104; or direct equivalents thereof.
- 4. A process according to claim 2 or 3 in which the CD25 binding molecule comprises at least
- a) one immunoglobulin heavy chain or fragment thereof which comprises (i) a variable domain comprising in sequence the hypervariable regions CDR1, CDR2 and CDR3 and (ii) the constant part or fragment thereof of a human heavy chain; said CDR1 having the amino acid sequence Arg-Tyr-Trp-Met-His, said CDR2 having the amino acid sequence Ala-Ile-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly, and said CDR3 having the amino acid sequence Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe and
  - b) one immunoglobulin light chain or fragment thereof which comprises (i) a variable domain comprising in sequence the hypervariable regions CDR1', CDR2' and CDR3' and (ii) the constant part or fragment thereof of a human light chain; said CDR1' having the amino acid sequence Ser-Ala-Ser-Ser-Ile-Ser-Tyr-Met-Gln, said CDR2' having the amino acid sequence Asp-Thr-Ser-Lys-Leu-Ala-Ser, and said CDR3' having the amino acid sequence His-Gln-Arg-Ser-Ser-Tyr-Thr;
  - or direct equivalents thereof.
  - 5. A process according to claim 4 in which the CD25 binding molecule comprises at least
    - a) one heavy chain which comprises a variable domain having the amino acid sequence shown in Seq.
       Id. No. 1 starting with amino acid at position 1 and ending with amino acid at position 117 and the constant part of a human heavy chain; and
    - b) one light chain which comprises a variable domain having the amino acid sequence shown in Seq. Id. No. 2 starting with glutamic acid at position 1 and ending with glutamic acid at position 104 and the constant part of a human light chain; or
      - direct equivalents thereof.

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- 6. A process according to claim 4 or 5 in which, in the CD25 binding molecule, the constant part or fragment thereof of the human heavy chain is of the γ<sub>1</sub> type and the constant part or fragment thereof of the human light chain is of the κ type.
- 50 7. A process for producing a multi-chain CD25 binding molecule which comprises
  - A) culturing an organism which is transformed with
    - i) a DNA construct which encodes a heavy chain or fragment thereof and comprises
      - a) a first part which encodes a variable domain comprising alternatively framework and hypervariable regions, said hypervariable regions being in sequence CDR1, CDR2 and CDR3 the amino acid sequences of which are shown in Seq. Id. No. 1; this first part starting with a codon encoding the first amino acid of the variable domain and ending with a codon encoding the last amino acid of the variable domain, and
      - b) a second part which encodes a heavy chain constant part or fragment thereof which starts with a codon encoding the first amino acid of the constant part of the heavy chain and ends

with a codon encoding the last amino acid of the constant part or fragment thereof, followed by a non-sens codon:

and with

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- ii) a DNA construct which encodes a light chain or fragment thereof and comprises
  - a) a first part which encodes a variable domain comprising alternatively framework and hypervariable regions; said hypervariable regions being in sequence CDR1', CDR2' and CDR3', the amino acid sequences of which are shown in Seq. Id. No. 2; this first part starting with a codon encoding the first amino acid of the variable domain and ending with a codon encoding the last amino acid of the variable domain, and
  - b) a second part which encodes a light chain constant part or fragment thereof which starts with a codon encoding the first amino acid of the constant part of the light chain and ends with a codon encoding the last amino acid of the constant part or fragment thereof followed by a nonsense codon; and
- B) recovering an active CD25 binding molecule from the culture.
- 8. A process for the preparation of an immunosuppressive composition comprising the step of mixing together at least one CD25 binding molecule defined in any one of claims 1 to 7 and at least one antigen binding molecule to at least one antigen other than CD25 which is characteristic of activated T-cells.
- 9. A process according to claim 8 which further comprises the admixture of cyclosporin A.
- 10. A process according to claim 8 or claim 9 in which the CD25 binding molecule is the chimeric CD25 antibody of Example 5 and the other antigen binding molecule is a chimeric CD7 antibody.
- 11. the use of a CD25 binding molecule defined in any one of claims 1 to 7 for the preparation of a medicament for immunosuppression of a mammalian system in conjunction with an antigen binding molecule recognising an antigen other than CD25 characteristic of activated T-cells.

## 30 Patentansprüche

## Patentansprüche für folgende Vertragsstaaten: AT, BE, CH, DE, DK, FR, GB, IT, LU, NL, SE

- CD25 bindendes Molekül, das mindestens eine Antigenbindungsstelle enthält, die mindestens eine Domäne enthält, die wiederum die hypervariablen Regionen CDR1, CDR2 und CDR3 in Folge enthält, wobei CDR1 die Aminosäuresequenz Arg-Tyr-Trp-Met-His, CDR2 die Aminosäuresequenz Ala-Ile-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly und CDR3 die Aminosäuresequenz Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe aufweist oder direkte Äquivalente hiervon.
- CD25 bindendes Molekül nach Anspruch 1, das mindestens eine Antigenbindungsstelle enthält, die folgendes umfaßt
  - a) eine erste Domäne, die in Folge die hypervariablen Regionen CDR1, CDR2 und CDR3 aufweist, wobei CDR1 die Aminosäuresequenz Arg-Tyr-Trp-Met-His, CDR2 die Aminosäuresequenz Ala-Ile-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly und CDR3 die Aminosäuresequenz Asp-Tyr-Gly-Tyr-Phe-Asp-Phe aufweist, und
  - b) eine zweite Domäne, die in Folge die hypervariablen Regionen CDR1', CDR2' und CDR3' aufweist, wobei CDR1' die Aminosäuresequenz Ser-Ala-Ser-Ser-Ile-Ser-Tyr-Met-Gln, CDR2' die Aminosäuresequenz Asp-Thr-Ser-Lys-Leu-Ala-Ser- und CDR3' die Aminosäueresequenz His-Gln-Arg-Ser-Ser-Tyr-Thr aufweist
- oder direkte Äquivalente hiervon.
  - 3. CD25 bindendes Molekūl nach Anspruch 1, das mindestens eine Antigenbindungsstelle enthält, die entweder eine Domäne mit der in Sequenzzuordnung Nr. 1 gezeigten Aminosäuresequenz beginnend mit der Aminosäure an Position 1 und mit der Aminosäure an Position 117 endend enthält oder eine wie oben beschriebene erste Domäne und eine zweite Domäne, die die in Sequenzzuordnung Nr. 2 gezeigte Aminosäuresequenz beginnend mit der Aminosäur an der Position 1 und mit der Aminosäure an der Position 104 endend enthält oder direkt Äquivalente hiervon.
  - 4. CD25 bindend s Molekül nach Anspruch 2 oder 3, das zumindest enthält

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a) eine schwere Kette eines Immunglobulins oder ein Fragment hiervon, die umfaßt (i) ein variable Domäne, die in Folg die hypervariablen Regionen CDR1, CDR2 und CDR3 enthält, und (ii) den konstanten Teil einer humanen schweren Kette oder ein Fragment hiervon, wobei CDR1 die Aminosäuresequenz Arg-Tyr-Trp-Met-His, CDR2 die Aminosäuresequenz Ala-Ile-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly und CDR3 die Aminosäuresequenz Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe aufweist, und

b) eine leichte Kette eines Immunglobulins oder ein Fragment hiervon, die umfaßt (i) eine variable Domäne, die in Folge die drei hypervariablen Regionen CDR1', CDR2' und CDR3' enthält, und (ii) den konstanten Teil einer humanen leichten Kette oder ein Fragment hiervon, wobei CDR1' die Aminosäuresequenz Ser-Ala-Ser-Ser-Ser-Ile-Ser-Tyr-Met-Gln, CDR2' die Aminosäuresequenz Asp-Thr-Ser-Lys-Leu-Ala-Ser- und CDR3' die Aminosäueresequenz His-Gln-Arg-Ser-Ser-Tyr-Thr aufweist, oder direkte Äquivalente hiervon.

- 5. CD25 bindendes Molekül nach Anspruch 4, das zumindest enthält a) eine schwere Kette, die eine variable Domäne mit der in der Sequenzzuordnung Nr. 1 gezeigten Aminosäuresequenz beginnend mit der Aminosäure an der Position 1 und mit der Aminosäure an der Position 117 endend und den konstanten Teil einer humanen schweren Kette enthält, und b) eine leichte Kette, die eine variable Domäne mit der in der Sequenzzuordnung Nr. 2 gezeigten Sequenz beginnend mit der Glutaminsäure an der Position 1 und mit der Glutaminsäure an der Position 104 endend und den konstanten Teil einer humanen leichten Kette enthält oder direkte Äquivalente hiervon.
- CD25 bindendes Molekül nach Anspruch 4 oder 5, worin der konstante Teil der humanen schweren Kette
  oder ein Fragment hiervon vom γ<sub>1</sub> Typ ist und der konstante Teil der humanen leichten Kette oder ein Fragment hiervon vom κ Typ ist.
  - DNA Molekül, das die Nukleotidsequenzen enthält, die in Folge die hypervariablen Regionen CDR1, CDR2 und CDR3, wie in der Sequenzzuordnung Nr. 1 gezeigt, kodieren.
- 8. DNA Konstrukt nach Anspruch 7, das eine schwere Kette oder ein Fragment hiervon kodiert und folgendes umfaßt
  - a) einen ersten Teil, der eine abwechselnd Framework und hypervariable Regionen enthaltende variable Domäne kodiert, wobei die in Folge auftretenden hypervariablen Regionen CDR1, CDR2 und CDR3 die in der Sequenzzuordnung Nr. 1 gezeigten Aminosäueresequenzen sind und dieser erste Teil mit einem Codon beginnt, das die erste Aminosäure der variablen Domäne kodiert und mit einem Codon endet, das die letzte Aminosäure der variablen Domäne kodiert, und
  - b) einen zweiten Teil, der einen konstanten Teil einer schweren Kette oder ein Fragment hiervon kodiert, der mit einem die erste Aminosäure des konstanten Teils kodierenden Codon beginnt und mit einem die letzte Aminosäure des konstanten Teils oder eines Fragments hiervon kodierenden Codon endet, gefolgt von einem Stopcodon.
  - 9. DNA Konstrukt nach Anspruch 8, das einen ersten Teil enthält, der eine variable Domäne mit der in der Sequenzzuordnung Nr. 1 gezeigten Aminosäuresequenz enthält, beginnend mit der Aminosäure an der Position 1 und mit der Aminosäure an der Position 117 endend oder direkte Äquivalente hiervon.
  - DNA Konstrukt nach Anspruch 7, das eine leichte Kette oder ein Fragment hiervon kodiert und folgendes umfaßt
    - a) einen ersten Teil, der eine abwechselnd Framework und hypervariable Regionen enthaltende variable Domäne kodiert, wobei die in Folge auftretenden hypervariablen Regionen CDR1', CDR2' und CDR3' die in der Sequenzzuordnung Nr. 2 gezeigten Aminosäuresequenzen sind und dieser erste Teil mit einem Codon beginnt, das die erste Aminosäure der variablen Domäne kodiert und mit einem Codon endet, das die letzte Aminosäure der variablen Domäne kodiert, und
    - b) einen zweiten Teil, der einen konstanten Teil einer leichten Kette oder ein Fragment hiervon kodiert, der mit einem die erste Aminosäure des konstanten Teils kodierenden Codon beginnt und mit einem die letzte Aminosäure des konstanten Teils oder eines Fragments hiervon kodierenden Codon endet, gefolgt von einem Stopcodon.
  - 11. DNA Konstrukt nach Anspruch 10, das einen ersten Teil enthält, der eine variable Domäne mit der in der Sequenzzuordnung Nr. 1 gezeigten Aminosäuresequenz enthält, beginnend mit der Aminosäure an der

Position 1 und mit der Aminosäure an der Position 104 endend oder direkt Äquivalent hiervon.

- 12. Verfahren zur Herstellung ein s mehrkettigen CD25 bindenden Moleküls, gek innzeichnet durch (i) Kultivieren eines Organismus, der mit einem DNA Konstrukt nach Anspruch 8 oder 9 und mit einem DNA Konstrukt nach Anspruch 10 oder 11 transformiert ist und (ii) Gewinnen eines aktiven CD25 bindenden Moleküls aus der Kultur.
- 13. Pharmazeutische Zusammensetzung zur Immunsuppression des humanen Immunsystems oder zur Behandlung von Malignitäten der CD25<sup>+</sup> Zellen oder zur Behandlung der HIV Infektion, die ein CD25 bindendes Molekül nach einem der Ansprüche 1 bis 6 und einen pharmazeutisch annehmbaren Träger oder ein pharmazeutisch annehmbares Verdünnungsmittel enthält.
- 14. Immunsuppressive Zusammensetzung, die eine Mischung von mindestens einem CD25 bindenden Molekül nach einem der Ansprüche 1 bis 6 und mindestens ein antigenbindendes Molekül gegenüber mindestens einem anderen Antigen als CD25, das für aktivierte T-Zellen charakteristisch ist, enthält.
  - 15. Zusammensetzung nach Anspruch 14, die zusätzlich Cyclosporin A enthält.
- Zusammensetzung nach Anspruch 14 oder 15, worin das antigenbindende Molekül anders als das CD
   bindende Molekül ein CD7 Antikörper ist.
  - 17. Zusammensetzung nach einem der Ansprüche 14 bis 16, worin die Antikörper chimäre Antikörper sind.
- 18. Doppelpackung, die getrennte Einheitsdosierungsformen von mindestens zwei Antikörpem gegenüber aktivierten T-Zellen enthält, wobei die Antikörper mindestens zwei verschiedene, für aktivierte T-Zellen charakteristische Antigene erkennen, wobei ein Antikörper ein CD25 bindendes Molekül nach einem der Ansprüche 1 bis 6 ist, zusammen mit Anleitungen zum Mischen oder zur gemeinsamen Verabreichung.
  - Doppelpackung nach Anspruch 18, die zusätzlich eine Einheitsdosierungsform eines immunsuppressiven Arzneimittels enthält.
  - 20. Verwendung eines CD25 bindenden Moleküls nach einem der Ansprüche 1 bis 6 in Kombination mit einem antigenbindenden Molekül, das ein anderes Antigen als CD25 erkennt, das für aktivierte T-Zellen charakteristisch ist, zur Herstellung eines Medikaments zur Immunsuppression des Säugetiersystems.

# 35 Patentansprüche für folgende Vertragsstaaten: ES, GR

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- 1. Verfahren zur Herstellung eines CD25 bindenden Moleküls, das mindestens eine Antigenbindungsstelle enthält, die mindestens eine Domäne enthält, die wiederum in Folge die hypervariablen Regionen CDR1, CDR2 und CDR3 enthält, wobei CDR1 die Aminosäuresequenz Arg-Tyr-Trp-Met-His, CDR2 die Aminosäuresequenz Ala-Ile-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly und CDR3 die Aminosäuresequenz Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe aufweist oder direkte Äquivalente hiervon, wobei dieses Verfahren gekennzeichnet ist durch einen Kultivierungsschritt einer Hybridomzellinie oder eines mit einem DNA Konstrukt transformierten Organismus, wobei das Konstrukt Nukleotidsequenzen enthält, die in Folge die hypervariablen Regionen CDR1, CDR2 und CDR3 enthalten, und Isolierung eines CD25 bindenden Moleküls aus der Kultur.
- Verfahren nach Anspruch 1, worin das CD25 bindende Molekül mindestens eine Antigenbindungsstelle enthält, die folgendes umfaßt
  - a) eine erste Domäne, die in Folge die hypervariablen Regionen CDR1, CDR2 und CDR3 aufweist, wobei CDR1 die Aminosäuresequenz Arg-Tyr-Trp-Met-His, CDR2 die Aminosäuresequenz Ala-Ile-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly und CDR3 die Aminosäuresequenz Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe aufweist, und
  - b) eine zweite Domäne, die in Folge die hypervariablen Regionen CDR1', CDR2' und CDR3' aufweist, wobei CDR1' die Aminosäuresequenz Ser-Ala-Ser-Ser-Ile-Ser-Tyr-Met-Gln, CDR2' die Aminosäuresequenz Asp-Thr-Ser-Lys-Leu-Ala-Ser- und CDR3' die Aminosäueresequenz His-Gln-Arg-Ser-Ser-Tyr-Thr aufweist

oder direkte Äquivalente hiervon.

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- 3. Verfahren nach Anspruch 1, worin das CD25 bindende Molekül mindestens eine Antigenbindungsstelle enthält, die entweder eine Domäne mit der in Sequenzzuordnung Nr. 1 gezeigten Aminosäuresequenz beginnend mit der Aminosäure an Position 1 und mit der Aminosäure an Position 117 end nd enthält, oder eine wie oben beschriebene erste Domäne und eine zweite Domäne, die die in Sequenzzuordnung Nr. 2 gezeigte Aminosäuresequenz beginnend mit der Aminosäure an der Position 1 und mit der Aminosäure an der Position 104 endend enthält oder direkte Äquivalente hiervon.
- 4. Verfahren nach Anspruch 2 oder 3, worin das CD25 bindende Molekül zumindest enthält a) eine schwere Kette eines Immunglobulins oder ein Fragment hiervon, die umfaßt (i) eine variable Domäne, die in Folge die hypervariablen Regionen CDR1, CDR2 und CDR3 enthält, und (ii) den konstanten Teil einer humanen schweren Kette oder ein Fragment hiervon, wobei CDR1 die Aminosäuresequenz Arg-Tyr-Trp-Met-His, CDR2 die Aminosäuresequenz Ala-Ile-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly und CDR3 die Aminosäuresequenz Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe aufweist, und
  - b) eine leichte Kette eines Immunglobulins oder ein Fragment hiervon, die umfaßt (i) eine variable Domäne, die in Folge die drei hypervariablen Regionen CDR1', CDR2' und CDR3' enthält, und (ii) den konstanten Teil einer humanen leichten Kette oder ein Fragment hiervon, wobei CDR1' die Aminosäuresequenz Ser-Ala-Ser-Ser-Ser-Ile-Ser-Tyr-Met-Gln, CDR2' die Aminosäuresequenz Asp-Thr-Ser-Lys-Leu-Ala-Ser- und CDR3' die Aminosäueresequenz His-Gln-Arg-Ser-Ser-Tyr-Thr aufweist, oder direkte Äquivalente hiervon.
  - 5. Verfahren nach Anspruch 4, worin das CD25 bindende Molekül zumindest enthält a) eine schwere Kette, die eine variable Domäne mit der in der Sequenzzuordnung Nr. 1 gezeigten Aminosäuresequenz beginnend mit der Aminosäure an der Position 1 und mit der Aminosäure an der Position 117 endend und den konstanten Teil einer humanen schweren Kette enthält, und b) eine leichte Kette, die eine variable Domäne mit der in der Sequenzzuordnung Nr. 2 gezeigten Sequenz beginnend mit der Glutaminsäure an der Position 1 und mit der Glutaminsäure an der Position 104 endend und den konstanten Teil einer humanen leichten Kette enthält, oder direkte Äquivalente hiervon.
    - 6. Verfahren nach Anspruch 4 oder 5, worin im CD25 bindenden Molekül der konstante Teil der humanen schweren Kette oder ein Fragment hiervon vom γ<sub>1</sub> Typ ist und der konstante Teil der humanen leichten Kette oder ein Fragment hiervon vom κ Typ ist.
- Verfahren zur Herstellung eines mehrkettigen CD25 bindenden Moleküls, gekennzeichnet durch
   A) Kultivierung eines Organismus, der transformiert ist mit
  - i) einem DNA Konstrukt, das die schwere Kette oder ein Fragment hiervon kodiert und folgendes enthält
    - a) einen ersten Teil, der eine abwechselnd Framework und hypervariable Regionen enthaltende variable Domäne kodiert, wobei die in Folge auftretenden hypervariablen Regionen CDR1, CDR2 und CDR3 die in der Sequenzzuordnung Nr. 1 gezeigten Aminosäueresequenzen sind und dieser erste Teil mit einem Codon beginnt, das die erste Aminosäure der variablen Domäne kodiert und mit einem Codon endet, das die letzte Aminosäure der vanablen Domäne kodiert, und
  - b) einen zweiten Teil, der einen konstanten Teil einer schweren Kette oder ein Fragment hiervon kodiert, der mit einem die erste Aminosäure des konstanten Teils kodierenden Codon beginnt und mit einem die letzte Aminosäure des konstanten Teils oder eines Fragments hiervon kodierenden Codon endet, gefolgt von einem Stopcodon, und mit
  - ii) einem DNA Konstrukt, das die leichte Kette oder ein Fragment hiervon kodiert und folgendes umfaßt
    - a) einen ersten Teil, der eine abwechselnd Framework und hypervariable Regionen enthaltende variable Domäne kodiert, wobei die in Folge auftretenden hypervariablen Regionen CDR1', CDR2' und CDR3' die in der Sequenzzuordnung Nr. 2 gezeigten Aminosäuresequenzen sind und dieser erste Teil mit inem Codon beginnt, das di erste Aminosäure der variablen Domäne kodiert und mit einem Codon endet, das die letzte Aminosäure d r variablen Domäne kodiert,
    - b) einen zweiten Teil, der einen konstanten Teil einer leichten Kette oder ein Fragment hiervon

kodiert, der mit einem die erste Aminosäure des konstanten Teils kodierenden Codon beginnt und mit einem die letzte Aminosäure des konstanten Teils oder eines Fragments hiervon kodierenden Codon ndet, gefolgt von einem Stopcodon, und

- B) Gewinnen eines aktiven CD25 bindenden Moleküls aus der Kultur.
- 8. Verfahren zur Herstellung einer immunsuppressiven Zusammensetzung, gekennzeichnet durch einen Mischungsschritt von mindestens einem CD25 bindenden Molekül nach einem der Ansprüche 1 bis 7 und mindestens einem antigenbindenden Molekül gegenüber mindestens einem anderen Antigen als CD25, das für aktivierte T-Zellen charakteristisch ist.
- 9. Verfahren nach Anspruch 8, das zusätzlich Cyclosporin A als Beimischung enthält.
- 10. Verfahren nach Anspruch 8 oder 9, worin das CD25 bindende Molekül der chimäre CD25 Antikörper von Beispiel 5 ist und das andere antigenbindende Molekül ein chimärer CD7 Antikörper ist.
- 11. Verwendung eines CD25 bindenden Moleküls nach einem der Ansprüche 1 bis 7 in Kombination mit einem antigenbindenden Molekül, das ein anderes Antigen als CD25 erkennt, das für aktivierte T-Zellen charakteristisch ist, zur Herstellung eines Medikaments zur Immunsuppression des Säugetiersystems.

#### Revendications

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Revendications pour les Etats contractants suivants : AT, BE, CH, DE, DK, FR, GB, IT, LU, NL, SE

- 1. Une molécule se liant à CD25 et qui comprend au moins un site de liaison à l'antigène comprenant au moins un domaine qui comprend dans l'ordre les régions hypervariables CDR1, CDR2 et CDR3, ladite région CDR1 ayant la séquence d'amino-acides Arg-Tyr-Trp-Met-His, ladite région CDR2 ayant la séquence d'amino-acides Ala-lle-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly, et ladite région CDR3 ayant la séquence d'amino-acides Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe, ou ses équivalents directs.
- 2. Une molécule se liant à CD25 selon la revendication 1 et qui comprend au moins un site de liaison à l'antigène comprenant :
  - a) un premier domaine comprenant dans l'ordre les régions hypervariables CDR1, CDR2 et CDR3, ladite région CDR1 ayant la séquence d'amino-acides Arg-Tyr-Trp-Met-His, ladite région CDR2 ayant la séquence d'amino-acides Ala-Ile-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly, et ladite région CDR3 ayant la séquence d'amino-acides Asp-Tyr-Gly-Tyr-Phe-Asp-Phe, et
  - b) un second domaine comprenant dans l'ordre les régions hypervariables CDR1', CDR2' et CDR3', ladite région CDR1' ayant la séquence d'amino-acides Ser-Ala-Ser-Ser-Ile-Ser-Tyr-Met-GIn, ladite région CDR2' ayant la séquence d'amino-acides Asp-Thr-Ser-Lys-Leu-Ala-Ser, et ladite région CDR3' ayant la séquence d'amino-acides His-GIn-Arg-Ser-Ser-Tyr-Thr, ou ses équivalents directs.
- 45 3. Une molécule se liant à CD25 selon la revendication 1 et qui comprend au moins un site de liaison à l'antigène comprenant soit un domaine ayant la séquence d'amino-acides indiquée dans la description de séquence Seq. Id. No 1 commençant par l'amino-acide en position 1 et se terminant par l'amino-acide en position 117, soit un premier domaine comme décrit ci-dessus et un second domaine ayant la séquence d'amino-acides indiquée dans la description de séquence Seq. Id. No 2, commençant par l'amino-acide en position 1 et se terminant par l'amino-acide en position 104, ou ses équivalents directs.
- 4. Une molécule se liant à CD25 selon la revendication 2 ou 3 et qui comprend au moins a) une chaîne lourde d'immunoglobuline ou l'un de ses fragments, qui comprend (i) un domaine variable comprenant dans l'ordre les régions hypervariables CDR1, CDR2 et CDR3 et (ii) la partie constante d'une chaîne lourde humaine ou l'un de ses fragments, ladite région CDR1 ayant la séquence d'amino-acides Arg-Tyr-Trp-Met-His, ladite région CDR2 ayant la séquence d'amino-acides Ala-lle-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly, et ladite région CDR3 ayant la séquence d'amino-acid s Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe, et b) une chaîne légère d'immunoglobuline ou l'un de ses fragments, qui comprend (i) un domaine variable

comprenant dans l'ordre les régions hypervariables CDR1', CDR2' et CDR3' et (ii) la partie constante d'une chaîne légère humaine ou l'un de ses fragments, ladite région CDR1' ayant la séquenc d'amino-acides Ser-Ala-Ser-Ser-Ile-Ser-Tyr-Met-Gln, ladite région CDR2' ayant la séquence d'amino-acides Asp-Thr-Ser-Lys-Leu-Ala-Ser, et ladite région CDR3' ayant la séquence d'amino-acides His-Gln-Arg-Ser-Ser-Tyr-Thr,

ou ses équivalents directs.

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- 5. Une molécule se liant à CD25 selon la revendication 4 et qui comprend au moins
  - a) une chaîne lourde qui comprend un domaine variable ayant la séquence d'amino-acides indiquée dans la description de séquence Seq. Id. No 1 commençant par l'amino-acide en position 1 et se terminant par l'amino-acide en position 117 et la partie constante d'une chaîne lourde humaine, et b) une chaîne légère qui comprend un domaine variable ayant la séquence d'amino-acides indiquée dans la description de séquence Seq. Id. No 2 commençant par l'acide glutamique en position 1 et se terminant par l'acide glutamique en position 104 et la partie constante d'une chaîne légère humaine, ou ses équivalents directs.
- 6. Une molécule se liant à CD25 selon la revendication 4 ou 5 et dans laquelle la partie constante de la chaîne lourde humaine ou l'un de ses fragments est du type γ₁ et la partie constante de la chaîne légère humaine ou l'un de ses fragments est du type κ.
- Une molécule d'ADN qui comprend la séquence de nucléotides codant dans l'ordre les régions hypervariables CDR1, CDR2 et CDR3 comme indiqué dans la description de séquence Seq. Id. No 1.
- 8. Une construction d'ADN selon la revendication 7 qui code une chaîne lourde ou l'un de ses fragments et qui comprend
  - a) une première partie qui code un domaine variable comprenant alternativement des régions de l'ossature et des régions hypervariables, lesdites régions hypervariables étant dans l'ordre CDR1, CDR2 et CDR3 dont les séquences d'amino-acides sont indiquées dans la description de séquence Seq. Id. No 1, cette première partie commençant par un codon qui code le premier amino-acide du domaine variable et se terminant par un codon qui code le dernier amino-acide du domaine variable, et b) une seconde partie qui code une partie constante de chaîne lourde ou l'un de ses fragments et qui commence par un codon codant le premier amino-acide de la partie constante de la chaîne lourde et se termine par un codon codant le dernier amino-acide de la partie constante ou de l'un de ses fragments, suivi par un codon non-sens.
- 9. Une construction d'ADN selon la revendication 8 qui comprend une première partie qui code un domaine variable ayant la séquence d'amino-acides indiquée dans la description de séquence Seq. Id. No 1 commençant par l'amino-acide en position 1 et se terminant par l'amino-acide en position 117, ou ses équivalents directs.
- 10. Une construction d'ADN selon la revendication 7 qui code une chaîne légère ou l'un de ses fragments et qui comprend
  - a) une première partie qui code un domaine variable comprenant alternativement des régions de l'ossature et des régions hypervariables, lesdites régions hypervariables étant dans l'ordre CDR1', CDR2' et CDR3' dont les séquences d'amino-acides sont indiquées dans la description de séquence Seq. Id. No 2, cette première partie commençant par un codon qui code le premier amino-acide du domaine variable et se terminant par un codon qui code le dernier amino-acide du domaine variable, et b) une seconde partie qui code une partie constante de chaîne légère ou l'un de ses fragments et qui commence par un codon codant le premier amino-acide de la partie constante de la chaîne légère et se termine par un codon codant le dernier amino-acide de la partie constante ou de l'un de ses fragments, suivi par un codon non-sens.
- 11. Une construction d'ADN selon la revendication 10, qui comprend une première partie qui code un domaine variable ayant la séquence d'amino-acides indiquée dans la description de séquence Seq. Id. No 2 commençant par l'amino-acide en position 1 et se terminant par l'amino-acide en position 104, ou ses équivalents directs.
- 12. Un procédé de préparation d'une molécule multi-chaîne se liant à CD25, qui comprend (i) la mise en culture d'un organisme qui est transformé avec une construction d'ADN selon la revendication 8 ou 9 et avec

une construction d'ADN selon la revendication 10 ou 11 et (ii) la récupération, à partir de la culture, d'une molécule active se liant à CD25.

- 13. Une composition pharmaceutique pour l'immunosuppression du système immunitaire humain, ou pour le traitement des tumeurs malignes de cellules CD25+ ou pour le traitement de l'infection au VIH, qui comprend une molécule se liant à CD25 selon l'une quelconque des revendications 1 à 6, et un support ou diluant pharmaceutiquement acceptable.
- 14. Une composition immunosuppressive comprenant un mélange d'au moins une molécule se liant à CD25 selon l'une quelconque des revendications 1 à 6, et d'au moins une molécule se liant à au moins un antigène, autre que CD25, qui est caractéristique des cellules T activées.
  - 15. Une composition selon la revendication 14 qui comprend en outre de la cyclosporine A.
- 15 16. Une composition selon la revendication 14 ou 15 dans laquelle la molécule se liant à un antigène, autre que la molécule se liant à CD25, est un anticorps CD7.
  - 17. Une composition selon l'une quelconque des revendications 14 à 16, dans laquelle les anticorps sont des anticorps chimères.
  - 18. Un emballage jumelé contenant des doses unitaires séparées d'au moins deux anticorps aux cellules T activées, lesdits anticorps reconnaissant au moins deux antigènes différents caractéristiques des cellules T activées, l'un des anticorps étant une molécule se liant à CD25 selon l'une quelconque des revendications 1 à 6, ensemble avec les instructions pour le mélange ou l'administration concomittante.
  - Un emballage jurnelé selon la revendication 18, qui comprend en outre une dose unitaire d'un médicament immunosuppresseur.
- 20. L'utilisation d'une molécule se liant à CD25 selon l'une quelconque des revendications 1 à 6 pour la préparation d'un médicament destiné à l'immunosuppression du système immunitaire chez les mammifères conjointement à une molécule se liant à un antigène reconnaissant un antigène, autre que CD25, caractéristique des cellules T activées.

## Revendications pour les Etats contractants sulvants : ES, GR

- Un procédé de préparation d'une molécule se liant à CD25 et qui comprend au moins un site de liaison à l'antigène comprenant au moins un domaine qui comprend dans l'ordre les régions hypervariables CDR1, CDR2 et CDR3, ladite région CDR1 ayant la séquence d'amino-acides Arg-Tyr-Trp-Met-His, ladite région CDR2 ayant la séquence d'amino-acides Ala-lle-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly, et ladite région CDR3 ayant la séquence d'amino-acides Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe, ou de ses équivalents directs, ledit procédé comprenant l'étape de mise en culture d'une lignée cellulaire d'hybridomes ou d'un organisme transformé avec une construction d'ADN comprenant des séquences de nucléotides codant dans l'ordre lesdites régions hypervariables CDR1, CDR2 et CDR3, et l'isolement, à partir de la culture, d'une molécule se liant à CD25.
- 2. Un procédé selon la revendication 1, dans lequel la molécule se liant à CD25 comprend au moins un site de liaison à l'antigène comprenant :
  - a) un premier domaine comprenant dans l'ordre les régions hypervariables CDR1, CDR2 et CDR3, ladite région CDR1 ayant la séquence d'amino-acides Arg-Tyr-Trp-Met-His, ladite région CDR2 ayant la séquence d'amino-acides Ala-Ile-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly, et ladite région CDR3 ayant la séquence d'amino-acides Asp-Tyr-Gly-Tyr-Phe-Asp-Phe, et
  - b) un second domaine comprenant dans l'ordre les régions hypervariables CDR1', CDR2' et CDR3', ladite région CDR1' ayant la séquence d'amino-acides Ser-Ala-Ser-Ser-Ile-Ser-Tyr-Met-GIn, ladite région CDR2' ayant la séquence d'amino-acides Asp-Thr-Ser-Lys-Leu-Ala-Ser, et ladite région CDR3' ayant la séquence d'amino-acides His-GIn-Arg-Ser-Ser-Tyr-Thr,
- ou ses équivalents directs.

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 Un procédé selon la revendication 1, dans l' quel la molécule se liant à CD25 comprend au moins un site de liaison à l'antigène comprenant soit un domain ayant la séquence d'amino-acides indiquée dans la

description de séquence Seq. Id. No 1 commençant par l'amino-acide en position 1 et se terminant par l'amino-acide en position 117, soit un premier domaine comme décrit ci-dessus et un s cond domaine ayant la séquence d'amino-acides indiquée dans la description de séquence Seq. Id. No 2, commençant par l'amino-acide en position 1 et se terminant par l'amino-acide en position 104, ou ses équivalents directs

- 4. Un procédé selon la revendication 2 ou 3, dans lequel la molécule se liant à CD25 comprend au moins a) une chaîne lourde d'immunoglobuline ou l'un de ses fragments, qui comprend (i) un domaine variable comprenant dans l'ordre les régions hypervariables CDR1, CDR2 et CDR3 et (ii) la partie constante d'une chaîne lourde humaine ou l'un de ses fragments, ladite région CDR1 ayant la séquence d'amino-acides Arg-Tyr-Trp-Met-His, ladite séquence CDR2 ayant la séquence d'amino-acides Ala-lle-Tyr-Pro-Gly-Asn-Ser-Asp-Thr-Ser-Tyr-Asn-Gln-Lys-Phe-Glu-Gly, et ladite région CDR3 ayant la séquence d'amino-acides Asp-Tyr-Gly-Tyr-Tyr-Phe-Asp-Phe, et
  - b) une chaîne légère d'immunoglobuline ou l'un de ses fragments, qui comprend (i) un domaine variable comprenant dans l'ordre les régions hypervariables CDR1', CDR2' et CDR3' et (ii) la partie constante d'une chaîne légère humaine ou l'un de ses fragments, ladite région CDR1' ayant la séquence d'amino-acides Ser-Ala-Ser-Ser-Ile-Ser-Tyr-Met-Gln, ladite région CDR2' ayant la séquence d'amino-acides Asp-Thr-Ser-Lys-Leu-Ala-Ser, et ladite région CDR3' ayant la séquence d'amino-acides His-Gln-Arg-Ser-Ser-Tyr-Thr,

ou ses équivalents directs.

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- 5. Un procédé selon la revendication 4, dans lequel la molécule se liant à CD25 comprend au moins a) une chaîne lourde qui comprend un domaine variable ayant la séquence d'amino-acides indiquée dans la description de séquence Seq. Id. No 1 commençant par l'amino-acide en position 1 et se terminant par l'amino-acide en position 117 et la partie constante d'une chaîne lourde humaine, et b) une chaîne légère qui comprend un domaine variable ayant la séquence d'amino-acides indiquée dans la description de séquence Seq. Id. No 2 commençant par l'acide glutamique en position 1 et se terminant par l'acide glutamique en position 104 et la partie constante d'une chaîne légère humaine, ou ses équivalents directs.
- 6. Un procédé selon la revendication 4 ou 5, caractérisé par le fait que dans la molécule se liant à CD25, la partie constante de la chaîne lourde humaine ou l'un de ses fragments est du type γ<sub>1</sub> et la partie constante de la chaîne légère humaine ou l'un de ses fragments est du type κ.
- Un procédé de préparation d'une molécule multi-chaîne se liant à CD25, qui comprend
   A) la mise en culture d'un organisme qui est transformé avec
  - i) une construction d'ADN qui code une chaîne lourde ou l'un de ses fragments et qui comprend a) une première partie qui code un domaine variable comprenant alternativement des régions de l'ossature et des régions hypervariables, lesdites régions hypervariables étant dans l'ordre CDR1, CDR2 et CDR3 dont les séquences d'amino-acides sont indiquées dans la description de séquence Seq. Id. No 1, cette première partie commençant par un codon qui code le premier amino-acide du domaine variable et se terminant par un codon qui code le dernier amino-acide du domaine variable, et
    - b) une seconde partie qui code une partie constante de chaîne lourde ou l'un de ses fragments et qui commence par un codon codant le premier amino-acide de la partie constante de la chaîne lourde et se termine par un codon codant le dernier amino-acide de la partie constante ou de l'un de ses fragments, suivi par un codon non-sens,

et avec

- ii) une construction d'ADN qui code une chaîne légère ou l'un de ses fragments et qui comprend a) une première partie qui code un domaine variable comprenant alternativement des régions de l'ossature et des régions hypervariables, lesdites régions hypervariables étant dans l'ordre CDR1', CDR2' et CDR3' dont les séquences d'amino-acides sont indiquées dans la description de séquence Seq. Id. No 2, cette première partie commençant par un codon qui code le premier amino-acide du domaine variable et se terminant par un codon qui code le dernier amino-acide du domaine variable, et
  - b) une second partie qui code une partie constante de chaîne légère ou l'un d ses fragments t qui commence par un codon codant le premier amino-acide d la partie constante de la chaîne légère et se termine par un codon codant le dernier amino-acide de la partie constante ou de

l'un de ses fragments, suivi par un codon non-sens, et B) la récupération, à partir de la culture, d'une molécule active se liant à CD25.

- 8. Un procédé de préparation d'une composition immunosuppressive, comprenant l'étape de mélanger en-5 semble au moins une molécule se liant à CD25 définie à l'une quelconque des revendications 1 à 7, et au moins une molécule se liant à au moins un antigène, autre que CD25, qui est caractéristique des cellules T activées.
- Un procédé selon la revendication 8, qui comprend l'admixtion de cyclosporine A.

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- 10. Un procédé selon la revendication 8 ou 9, dans lequel la molécule se liant à CD25 est l'anticorps CD25 chimère de l'exemple 5 et l'autre molécule se liant à un antigène est un anticorps CD7 chimère.
- 11. L'utilisation d'une molécule se liant à CD25 définie à l'une quelconque des revendications 1 à 7, pour la 15 préparation d'un médicament destiné à l'immunosuppression du système immunitaire chez les mammifères conjointement à une molécule se liant à un antigène et reconnaissant un antigène, autre que CD25, caractéristique des cellules T activées.

Fig. 1A

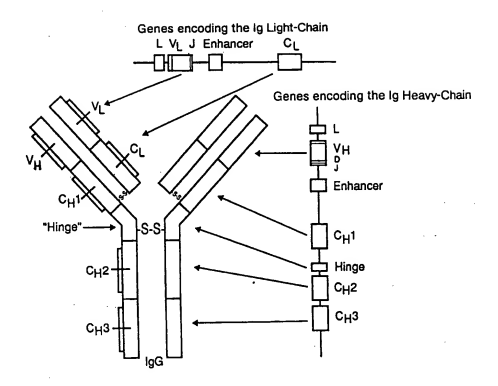
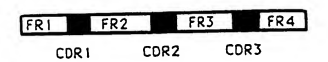


Fig. 1B



# Fig. 2A

(kb)

22 ---

9.0 ---

6.5 — 2 5 5

5.0 ---

4.3 ——

3.5 —

12 34

# Fig. 2B

[kb]

6.5 ----

5.0 ---

4.3 -

35-43 2 I

Fig. 3A

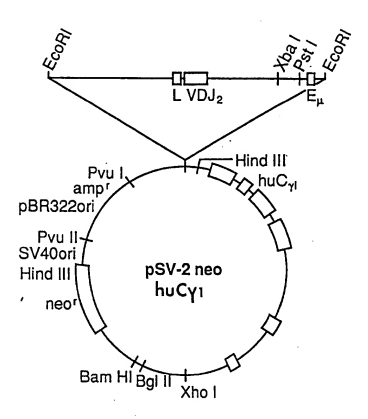
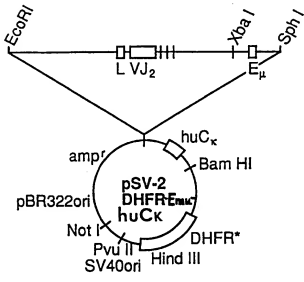
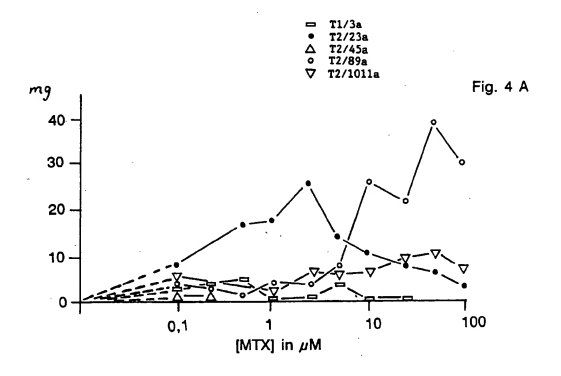
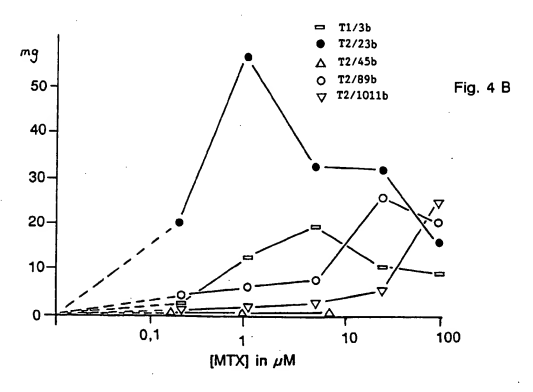


Fig. 3B







# Figure 5

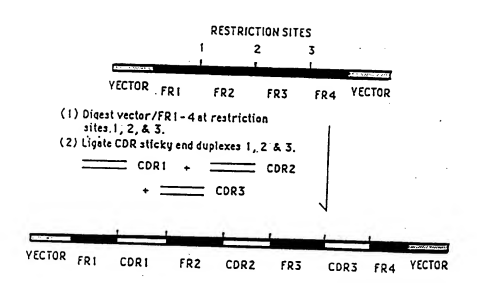


Fig. 6

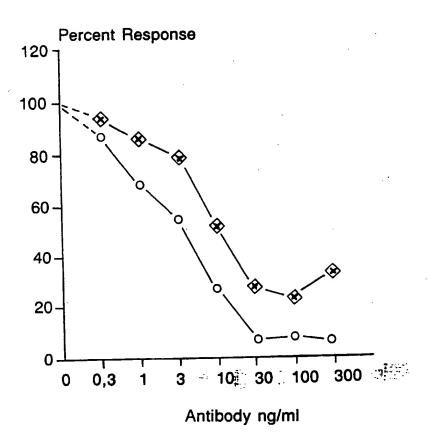


Fig. 7

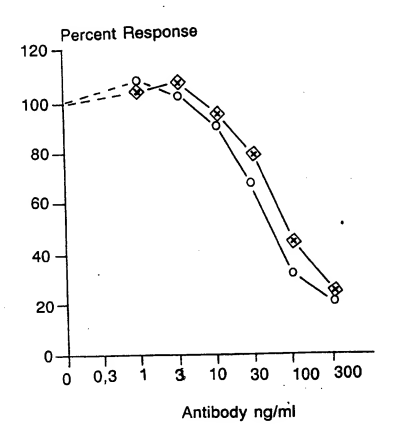
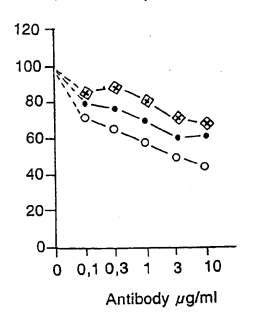


Fig. 8 A

- Fig. 8 B

# Percent Response



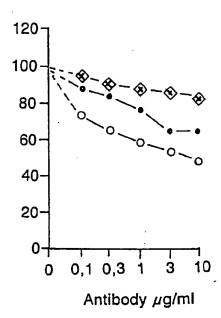


Fig. 8 C

Fig. 8 D



